PALEOGROUNDWATER IN THE MOUTERE GRAVEL AQUIFERS NEAR NELSON, NEW ZEALAND

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ABSTRACT. Radiocarbon, ¹⁸O, and chemical concentrations have been used to identify groundwater recharged during the last ice age near Nelson, New Zealand. Moutere Gravel underlies most of the Moutere Depression, a 30-km-wide system of valleys filled with Plio-Pleistocene gravel. The depression extends northwards into Tasman Bay, which was above sea level when the North and South Islands of New Zealand were connected during the last glaciation. The aquifers are tapped by bores up to 500 m deep. Shallow bores (50–100 m) tap "pre-industrial" Holocene water (termed the "modern" component) with ¹⁴C concentrations of 90 ± 10 percent modern carbon (pMC) and δ^{18} O values of $-6.8 \pm 0.4\%$, as expected for present-day precipitation. Deeper bores discharge water with lower ¹⁴C concentrations and more negative ¹⁸O values resulting from input of much older water from depth. The deep end-member of the mixing trend is identified as paleowater (termed the "glacial" component) with ¹⁴C concentration close to 0 pMC and more negative ¹⁸O values (-7.6%). Mixing of the modern and glacial components gives rise to the variations observed in the ¹⁴C, ¹⁸O, and chemical concentrations of the waters. Identification of the deep groundwater as glacial water suggests that there may be a large body of such water onshore and offshore at deep levels. More generally, the influence of changing sea levels in the recent past (geologically speaking) on the disposition of groundwaters in coastal areas of New Zealand may have been far greater than we have previously realized.

INTRODUCTION

Groundwater in the Moutere Valley is an important resource for horticulture. Deep bores revealed the hitherto unknown water resource in the early 1980s, after shallow bores were found to provide limited and unreliable supplies. The present work uses isotope and chemical measurements in conjunction with other data (Thomas 1989) to improve understanding of the system (Stewart and Thomas 2002).

The term "paleowater" refers to groundwater that can be clearly identified by means of its radiocarbon date, and/or by another isotopic or noble gas signature, as originating in the colder climatic conditions of the late Pleistocene (Edmunds 2001). The objectives of this paper are to establish the deep Moutere Gravel groundwater as "paleo" water, and to consider some implications for New Zealand groundwaters.

Hydrogeology

The Moutere Depression is a 30-km-wide system of valleys between the Tasman Mountains and the ranges of east Nelson at the top of the South Island (Figure 1a). Voluminous Plio-Pleistocene gravels (including Moutere Gravel) are preserved in the depression and have been incised by the Motueka, Moutere, and Waimea Rivers (Rattenbury et al. 1998). Geophysical interpretation of seismic data and petroleum bores indicate that the depression reaches depths of 2500 m on the eastern side (Lihou 1992). The depression formed in the Pliocene-Pleistocene during uplift of the Tasman Mountains and the east Nelson Ranges.

Moutere Gravel is a uniform yellow-brown, clay-bound gravel, with deeply weathered clasts almost entirely of Torlesse-derived sandstone and semi-schist. Well-rounded, quartzofeldspathic sandstone clasts in a brown weathered muddy sand matrix comprise the bulk of the gravel.

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Figure 1 (a) Geology of the Moutere Depression, northwest Nelson. The contact with Separation Point Granite (ks) defines the western boundary, while the eastern boundary is against the upward-faulted east Nelson Ranges. (b) Geological setting of the Moutere Valley showing bore locations. Fine lines show the surface geology comprising Separation Point Granite (ks) to the northwest and Moutere Gravel (units tm_1 and tm_2) to the southeast. The valley floor contains reworked Moutere Gravel (uk₃).

The Moutere Valley lies south of Motueka (Figure 1b). Most of the Moutere River catchment comprises Moutere Gravel (marked tm). The catchment is underlain by a north-east trending basinal structure in which Moutere Gravel reaches a maximum depth of about 600 m (Figure 2). A basement high running through Ruby Bay forms the southern flank of the basin. Basement granite then descends south of Ruby Bay to the Waimea Fault on the east side of the Waimea Plains reaching 2500 m. Groundwater in Moutere Gravel south of the Ruby Bay high is largely unexplored.

The floor of the Moutere Valley has a Quaternary infilling (uk) which derives from reworked Moutere Gravel (Figures 1b, 2). Shallow bores (<20 m) draw limited and often unreliable supplies of water from the valley infill, showing that permeabilities are low. Shallow bores on the Moutere Hills also tend to have low yields and unreliable supplies. Rainfall is the main recharge source of these supplies, with possibly variable contributions from local streams (Thomas 1989).

The hydrogeology has been described by Thomas (1989, 1991, 1992, 2001). Significant groundwater resources are found in deep Moutere Gravel aquifers. The 3 aquifers in vertical succession are the Shallow Moutere Aquifer (SMA), Middle Moutere Aquifer (MMA), and Deep Moutere Aquifer (DMA) (Figure 2). Intervening leaky clay layers containing carbonaceous material constrain the aquifers. Groundwater heads are generally higher and yields improve significantly with depth. Many deep bores (>50 m) are artesian. The deep bores are cased to 30–50 m depth and then are generally screened or open to the bottom, so that they are likely to draw on water from several depths. Thomas (1989) has identified feed zones in several bores from downhole measurements.



Figure 2 Cross-section showing the structure of the basin underlying the Moutere Catchment. The trace of the cross-section is parallel to the fault in Figure 1b. Inferred recharge patterns for the Moutere aquifers are shown (Stewart and Thomas 2002).

The Moutere groundwater resource north of the Ruby Bay basement high is divided into 2 zones (the western and eastern zones) by a major fault (Figure 1b). Pressure communication between bores is limited across this barrier. These zones are used for management of the resource (Thomas 2001). A third groundwater zone (southern) is defined south of the Ruby Bay high for discussion purposes.

Recharge to the aquifers is believed to occur by infiltration of rainfall where the Moutere Gravels outcrop. Two units have been identified by geological mapping, a lower unit (tm_1) and an upper unit (tm_2). The lower unit outcrops only in the southwest of the area (Figures 1b and 2), dips gently northeast into the Moutere Valley, and correlates with the DMA and MMA. The upper unit correlates with the SMA. Thomas (1989, 1992) proposed a recharge model in which rainfall infiltrated only where the lower unit outcrops in the Rosedale Hills area. This was revised to include recharge from both outcropping units of Moutere Gravel (Figure 2, Stewart and Thomas 2002). However, such recharge only applies to the "modern" component because conditions would have been very different during recharge of the "glacial" component. And no modern recharge (i.e. tritium or CFC-bearing water) has yet been observed in the bores within the valley, despite groundwater levels generally recovering from their summer drawdown during the subsequent winter and spring.

SAMPLING AND METHODS

Sampling

Groundwater bores were purged of at least 3 casing volumes before samples were taken. Bottles were flushed with the water to be sampled, emptied, refilled with water and allowed to overflow, then carefully sealed to prevent evaporation. Samples of water were collected in 28-mL glass bottles for ¹⁸O, 500-mL bottles for carbon isotopes, and 1.1-L bottles for tritium. The carbon isotope samples were collected air-free. Bore locations are shown in Figure 1b. (More details on the bores are given in Stewart and Thomas 2002.)

Rafter Radiocarbon Seawater Line Method

The 150-mL samples were extracted using the Rafter Radiocarbon standard seawater method. A seawater vessel is prepared before each extraction. Four mL of ortho-phosphoric acid is added to the

side arm of the flask, and a magnetic stirrer and anti-bumping granules added to the bottom of the flask. The flask and contents are "tared" on a 2 decimal place balance. The 150-mL portion of water to be processed was removed using a plastic syringe from each groundwater sample. The water was transferred to the bottom of the glass seawater vessel using the syringe. The water was added carefully so the acid and water did not mix. Care was taken not to expose the water samples to atmospheric air as much as possible, by flushing the vessel with N_2 gas through a thin plastic hose placed into the spout of the vessel during addition of the sample. The sample bottle was sealed and weighed. The sample and acid were shaken together for approximately 1 min to mix, and attached to the seawater vacuum line. Room air was pumped from the connection between the seawater vessel and the vacuum line to a desirable vacuum. Two dry ice ethanol dewars are placed on the first 2 traps to collect water vapor, and 2 liquid nitrogen dewars are placed on the following 2 traps to collect carbon dioxide. The water sample in the seawater vessel is stirred throughout the transfer with a magnetic stirrer to aid transfer of the gas. The water bubbles vigorously during the vacuum line has again reached the baseline vacuum, which takes approximately 20 min.

The carbon dioxide gas is vacuum-distilled twice to further dry and purify it before measurement of the pressure. The amount of CO₂ gas collected is measured in a known volume, and the amount of dissolved inorganic carbon in the sample calculated. The carbon dioxide gas is made into a graphite target for measurement of ¹⁴C by AMS. ¹⁴C concentration is expressed as percent modern carbon (pMC), where the activity of "modern carbon" is taken as 95% of the activity of the NBS oxalic acid standard in 1950. Errors depend on concentration, ranging from ±0.07 pMC for near zero to ±0.8 pMC for 100 pMC concentrations.

Part of the gas is retained for measurement of ¹³C by mass spectrometry. ¹³C concentrations are expressed as δ values with respect to VPDB (Vienna PDB, the international standard). The measurement errors are ±0.1‰.

¹⁸O and Tritium

For ¹⁸O measurement on the waters, 2 mL of the water is isotopically equilibrated with CO₂ gas at 29 °C for 2 hr, and then the CO₂ is analyzed in a stable isotope mass spectrometer (Hulston et al. 1981). ¹⁸O concentrations are given as δ values with respect to VSMOW (Vienna SMOW, the international standard). Measurement errors are ±0.1‰.

Tritium samples are distilled, enriched in tritium using electrolysis by a factor of about 80, and are then counted in a Quantalus low-background liquid scintillation counter for several weeks (Taylor 1994). Tritium concentrations are given as tritium units (TU), where 1 TU is 1×10^{-18} . Detection limits (errors for near-zero concentrations) are ± 0.003 TU.

Chemical Data

Water samples have been collected from groundwater bores in the Moutere Valley by Tasman District Council hydrologists for a number of years. Samples for cations were field filtered and acidified with high-purity nitric acid. Samples for anions were field filtered and kept below 4 °C until analyzed, and bicarbonate samples were collected unfiltered, kept below 4 °C, and analyzed within 48 hr of collection. Samples were analyzed by the Cawthron Institute in Nelson. Methods for cation analyses include Atomic Adsorption and ICP-OES, and for anion analyses include auto titrator, auto analyzer, and ion chromatography.

RESULTS

Carbon Isotope Compositions

The ¹³C concentrations in dissolved inorganic carbon (DIC) from Moutere are plotted against the reciprocal of the bicarbonate concentration (1/bicarbonate) in Figure 3a. The ¹³C values are remarkably uniform around the values -20 to -24% (except for 1 sample, see below). This range shows that all of the carbon is sourced from organic matter within the soil or aquifers. Waters gain dissolved CO₂ by plant respiration and oxidation of organic matter as they pass through the soil, where CO₂ partial pressures are commonly 10 to 100 times those in the atmosphere. A second source of CO₂ is from carbonaceous matter within the aquifers, in this case from the intervening clay layers between the aquifers. Oxidation of such matter produces CO₂ if chemical and/or microbiological conditions are suitable (bacteria are needed to catalyze redox reactions between water and organic matter). The dissolved CO₂ reacts with rocks to become neutralized to HCO₃⁻ and CO₃²⁻ ions (Clarke and Fritz 1997). Bicarbonate is the dominant form of carbonate in the present waters.



Figure 3 (a) δ^{13} C values versus (HCO₃)⁻¹ for bore waters; (b) ¹⁴C values versus (HCO₃)⁻¹ for bore waters. The line connects the glacial (zero ¹⁴C, 200 mg/L bicarbonate) and modern (90 pMC ¹⁴C, 125 mg/L bicarbonate) end-members.

Only 1 sample has ¹³C outside the range of -20 to -24‰, namely, 8107 with ¹³C of -13.3‰. This sample is likely to have reacted with marine carbonate, which has the effect of moving the sample towards the ¹³C value of such carbonate (approximately 0‰). The dotted line in Figure 3a shows the effect of reaction with marine carbonate. Carbon from carbonate rock would have zero ¹⁴C; hence, it would have a diluting effect on the ¹⁴C concentration. The short dotted line in Figure 3b shows the effect of reaction with carbonate on this sample.

Groundwater dating by ¹⁴C is complicated by changes in ¹⁴C activity in the atmosphere during the late Pleistocene and Holocene, and by dilution of ¹⁴C in groundwater by dead carbon derived from soils and rocks when carbon-bearing solutions penetrate underground. Most of the ¹⁴C in groundwater is gained from the soil, where CO₂ accumulates by root respiration and decay of vegetation, as noted above. The ¹⁴C in DIC is susceptible to reaction and dilution with dead carbon from carbonate, other minerals, and organic matter in the soil and groundwater zones. A dilution factor *q* is used to account for the resulting dilution of ¹⁴C (Clarke and Fritz 1997). The age equation is written

$$t = (1/\lambda) \cdot \ln(q \cdot a_o/a_t) \tag{1}$$

where a_o is the initial ¹⁴C activity ($q.a_o$ the diluted initial activity in the groundwater), a_t is the ¹⁴C activity in groundwater after time t (i.e. when measured), and λ is the ¹⁴C decay constant ($1/\lambda = T_{\frac{1}{2}}/\ln 2 = 8267$ yr). The apparent simplicity of this equation is deceptive. Numerous methods have been proposed for estimating q, based on the chemical and ¹³C composition of the groundwater (a summary of methods is given in Clarke and Fritz [1997]).

Figure 3b shows ¹⁴C concentrations plotted against $1/\text{HCO}_3$. The ¹⁴C concentrations show a wide range of values, from almost 0 to 100 pMC. Shallow bores have ¹⁴C concentrations in the range of 90 ± 10 pMC; this is taken as the initial ¹⁴C activity. (None of the waters have concentrations high enough to indicate the presence of ¹⁴C from nuclear weapons testing and hence are "pre-bomb"; i.e., they were all recharged before the early 1950s.)

The low ¹⁴C concentrations in the deep bores along with the low ¹³C values (which indicate absence of marine carbonates) show that the waters have had long residence times in the deep aquifer. None of the waters have zero ¹⁴C concentration (the lowest are 8.6 pMC [western and eastern zones] and 2.4 pMC [southern zone]), but the waters are mixtures because the bores are open to the aquifer from 50 m down to as much as 500 m depth. This allows water from several depth levels to contribute to the discharges. The ¹⁴C, ¹⁸O, and some chemical constituents show variations consistent with such mixing; Figure 4a shows the mixing line between ¹⁴C and $\delta^{18}O$.



Figure 4 (a) ¹⁸O versus ¹⁴C for bore waters; the older waters (low ¹⁴C) have more negative ¹⁸O values. (b) ¹⁸O values versus bore depths. A line is fitted to the points and connects deep (-7.6%, 600 m) and shallow (-6.8%, 0 m) end-members.

The data have been interpreted in terms of mixing between 2 types of water, a shallow component with ${}^{14}C$ concentration of $90 \pm 10\,$ pMC and a deep component with 0 pMC. The shallow component is designated "modern," but it has zero tritium and CFC concentrations and, therefore, is "pre-industrial" water with age in the hundreds of years (Stewart and Thomas 2002).

The deep component was recharged during the last glaciation and is the "glacial" component. A minimum mean age can be estimated using Equation 1. We assume that the ¹⁴C activity of the deep water is less than 1 pMC, and that of the shallow water is 90 pMC. The bicarbonate concentrations in the deep and shallow waters are 200 and 60 mg/L, respectively (Figure 5a); hence, the dilution factor (q) is 60/200. These give a mean age greater than 27,000 yr. This is probably an underestimate, but an age within the glacial period is what is important, not the precise age.



Figure 5 (a–d) 18 O versus HCO₃, Ca, Cl, and Na concentrations. Lines connect end-members at 18 O values of -7.6% (glacial water) and -6.8% (modern water).

¹⁸O Concentrations

 δ^{18} O values give information on the source and conditions of recharge to the groundwater because there is generally a relationship between the location of recharge and its ¹⁸O value (Stewart and Morgenstern 2001). ¹⁸O concentrations in water are expressed as δ values with respect to VSMOW (Vienna Standard Mean Ocean Water).

The mean δ^{18} O value of precipitation at Moutere is discussed in Stewart and Thomas (2002). They conclude that precipitation-recharged groundwater in the region under present-day conditions will have mean δ^{18} O values in the range of $-6.8 \pm 0.4\%$. This comes from analysis of monthly samples of precipitation from several nearby sites and from locally-fed springs and shallow bores.

Estimation of the mean ¹⁸O value of precipitation during the last glaciation is more problematical. The values will have been affected by at least 3 competing processes: 1) the sea would have been enriched in ¹⁸O because low-¹⁸O ice would have been locked up in ice sheets worldwide, leading to higher mean ¹⁸O values; 2) the atmospheric temperature would have been lower, leading to greater isotopic fractionation between atmospheric water vapor and precipitation, and, therefore, to lower mean ¹⁸O values; and 3) the sea would have been far from its present position (by up to several hundred km for Moutere), leading to a more continental climate and, therefore, to lower mean ¹⁸O values.

ues. Evidence suggests that glacial age groundwater at Moutere would have been more negative than late Holocene groundwater by about 1‰. (e.g., The δ^{18} O value of calcite deposited on stalactites near Hamilton, New Zealand, was 0.95‰ lower during the last glacial maximum than present deposits [Hendy and Wilson 1968]. This difference would have been mainly due to the δ^{18} O difference in the waters flowing over the stalactite at these times [Hendy and Wilson 1968].)

The ¹⁸O values of the groundwaters were found to be in the range of -6.4 to -7.5%, except for 1 bore in the southern zone with ¹⁸O of -8.3%. This is a wider range than expected for Holocene waters in the area. The ¹⁸O values also show relationships with bore depth (Figure 4b), with lower ¹⁸O values at depth, and with ¹⁴C as seen above. These observations support the idea of 2 types of water (modern and glacial). The ¹⁸O of the glacial component is found to be -7.6% from the mixing diagrams.

Hence, the favored explanation is that there are 2 types of water in the Moutere aquifers. The first is a body of "fossil" water, occupying deeper levels of the Moutere aquifers, which has lower ¹⁸O values because it was recharged during the last glacial period (in the Pleistocene). Above this is "modern" water, recharged in the last few hundred years of the Holocene, and with ¹⁸O values much the same as today. These 2 end-member compositions are shown by the large hollow squares in the figures; the line connecting them shows the effect of mixing of these waters. (Older Holocene water may also be present in the modern component.)

The southern zone bores do not draw on a consistent deep component, unlike the deep bores in the western and eastern zones, and show much more scatter. Part of the reason may be that deep permeabilities are not so high in the south and, consequently, the southern bores draw more water from higher levels in the aquifers than in the western and eastern zones.

Chemical Compositions

The chemical compositions of the western and eastern zone waters are affected by mixing of the 2 water types. The glacial water at depth appears to have a nearly uniform composition, whereas the modern water at shallow depths has a more varied composition. This section looks at data for 4 of the chemical constituents, which show patterns representative of many of the other chemical constituents (Stewart and Thomas 2002).

Bicarbonate and calcium (Figures 5a, b) are readily affected by reaction with soil and rock, and the deep component has higher concentrations of these constituents than the shallow component (the compositions of the end components are indicated by the large squares in the plots). The pattern is of increasing concentration with decreasing δ^{18} O value, which is indicative of a mixing process. This is consistent with the deep (glacial) component having had much greater interaction with the rock because of its much longer residence time.

Chloride concentrations in groundwater often reflect marine influence on rainfall (i.e. rainout of seasalt particles in the atmosphere) because it is often an unreactive solute underground (Edmunds 2001). For rainfall-recharged groundwater, evapotranspiration causes enrichment of chloride while passing through the soil. A third influence is seawater trapped within parts of the aquifer during past higher sea level stands, or present-day intrusion of seawater. The plot of chloride versus ¹⁸O (Figure 5c) shows glacial water (Cl ~ 5.5 mg/L, ¹⁸O ~ -7.6‰) mixing with modern water containing a range of chloride values (5.5 to 18 mg/L, ¹⁸O ~ -6.8‰). The graph has 2 limbs. Most of the western and eastern bores have chloride near 5.5 mg/L. However, bores on the seaward side of the Moutere Valley (8110, 8423, 8391, and 8089) plot on the upper limb with higher chloride concentrations. This is considered to reflect their location near the sea, where they receive rainfall that has higher chloride concentrations from Tasman Bay. If so, this shows that the recharge has occurred locally. The lower limb chloride concentration is very low (5.5 mg/L), and probably reflects a continental effect during both glacial and modern times due to the presence of the high Tasman Mountains west of Moutere, which lie at right angles to the prevailing westerlies. Hence, it appears that the input of chloride to groundwater in this region might have been much the same in glacial times as it was in pre-industrial (late Holocene) times.

Sodium (Figure 5d) has similarities to chloride but is more affected by interaction with aquifer rocks. Seawater influence is shown by higher sodium concentrations in 8050 and 8007 (not plotted). The plot with ¹⁸O shows 2 limbs as with chloride, but the upper limb values are more scattered (although it still includes 2 of the same bores, i.e., 8110 and 8089). Most of the samples lie on the lower limb, forming a trend towards higher sodium at depth because of increased water/rock interaction in the deep component.

The southern zone points do not generally conform to the patterns shown by the western and eastern zone points, although they are within the same range of values. As before, this is tentatively attributed to lower permeability at depth in the southern zone, leading to more input of waters from higher levels.

Tritium and CFCs

Tritium and CFC concentrations have been measured for a number of the bores (Stewart and Thomas 2002). The results show that the bore waters did not contain detectable tritium or CFC concentrations when first measured (as expected) and continue to show none. (If anything, ¹⁴C concentrations appear to be decreasing rather than increasing since 1988.) On the other hand, groundwater levels, which are drawn down during the summer, generally recover during the subsequent winter and early spring. Thus, recharge is occurring, but this new water (which is seen in shallow bores in the hilly Moutere Gravel outcrop areas) must be near the surface in the recharge areas and has not yet found its way to the aquifers tapped by the deep bores, or younger water is slowly penetrating into the deep aquifers, but dilution with old water has been sufficient to keep concentrations at less than detectable levels up to now.

DISCUSSION

¹⁴C and ¹⁸O concentrations give linear relationships with depth and with each other in the western and eastern zones, showing that 2 types of water are present. Shallow groundwater has ages of hundreds of years and δ^{18} O values like present-day rainfall (around –6.8‰). The deep component has low ¹⁴C concentration and ¹⁸O of –7.6‰. These are the modern and glacial water types identified above. The western and eastern bores discharge either modern or mixtures of modern and glacial water, none discharge pure glacial water. The glacial water is believed to have been recharged during part of the last glaciation (20–50,000 yr ago).

This is the first time that the δ^{18} O shift between modern and glacial groundwaters has been reported for a New Zealand system. The measured difference of -0.8% can be compared with differences of +0.5% (Portugal), -1.3% (France), -1.9% (England), and -2.5% (Poland), measured in coastal regions of Europe (from Figure 7 in Loosli et al. 2001).

The late Quaternary covers a period of strong climatic changes in the New Zealand landscape brought about by the last glaciation. So much ice was locked up in ice sheets that the sea level was about 130 m lower than at present at the last glacial maximum (LGM). This resulted in large

increases in the New Zealand land area during which time new hydrogeological systems were established offshore from the present coastline. Flow within deep aquifers would have been much more active because of the much greater heads due to lower sea level. The warmer climate during the Holocene saw the sea level rise rapidly to its present level as the ice sheets melted. This caused water tables in groundwater systems to rise and hydraulic heads to diminish, causing profound changes in flow regimes. In particular, flow at deep levels would have become much more sluggish. The present-day configuration of groundwaters in coastal areas is, therefore, to a large extent the result of circulation of freshwater to deeper levels in response to the lower sea level during the last glaciation (Edmunds 2001). The Holocene flow patterns have been superimposed on top of the glacial flow patterns of 10,000 yr and more ago.

Figure 6a shows the eustatic sea level curve for the last 140,000 yr (Edmunds et al. 2001, after Shackleton 1987). Until 7000 yr ago, the sea had not been at its present level for 100,000 yr; it was much lower for most of that time (especially for the last 50,000 yr of it). The minimum level was reached about 18,000 yr ago (at the LGM). Between 14,000 and 7000 yr ago, the sea level rose rapidly (with fluctuations), and in New Zealand is believed to have reached levels 2–3 m above the present level by 4000 yr ago (Stevens 1990). The level has since declined to the present level.



Figure 6 (a) Eustatic sea level curve for the past 140,000 yr; (b) New Zealand shoreline in the last glacial maximum (about 18,000 yr ago).

Figure 6b shows the inferred New Zealand shoreline during the LGM (Stevens 1990). The New Zealand land mass was doubled and the main islands of New Zealand were joined by a large plain. Moutere was an inland location, hundreds of km from the nearest sea. A considerable amount of the

former Pleistocene freshwater body (detected in this work) could now be present extending under Tasman Bay. It is probable that there is very little natural flow through the deep aquifer (i.e. little natural flow towards outlets under the sea). The lack of ¹⁴C indicates that no substantial Holocene recharge has taken place into this water body. Some of the water is evidently now being extracted by the deep Moutere bores, causing marked drawdowns in water levels during the late summer. However, levels recover during the winter. Hence, recharge is occurring and there must be a tendency for shallow water to penetrate deeper as the deep water is extracted.

Where repeated measurements have been made (bores 8404 and 8407), there appears to be a trend towards ages becoming older due to exploitation. This suggests that more of the deep water is being extracted because permeabilities at depth allow more efficient recovery (i.e. water flows more rapidly in the deep aquifer).

Ice age groundwaters have been observed in deep aquifers in other parts of New Zealand. Two of them are in basins in the North Island bordering Cook Strait (see Figure 7). Taylor and Evans (1999) reported a ¹⁴C age of 31,000 yr for groundwater at a depth of 238.4–241.4 m in the Whenuakura Formation near Hawera. Another deep groundwater at Shannon was reported as having a ¹⁴C age of 40,000 yr (CB Taylor, personal communication; see also Taylor et al. 2001). Both bores are in sedimentary formations.



Figure 7 Shoreline in the Cook Strait region during the last glacial maximum and ¹⁴C ages of some deep groundwaters.

These observations again suggest circulation of freshwater to deeper levels in response to the lower sea level during the last glaciation. The influence of changing sea levels in the recent past (geologically speaking) on the disposition of groundwaters in coastal areas of New Zealand may have been far greater than we have previously realized, and New Zealand may be surrounded by a "skirt" of pristine paleo-groundwaters at deep levels.

CONCLUSIONS

¹⁴C, ¹⁸O, and chemical concentrations have been used to identify groundwater in the Moutere Gravel aquifers that was recharged during the last ice age. The aquifers are tapped by bores up to 500 m deep. Shallow bores (50–100 m) tap pre-industrial Holocene water (termed the "modern" component) with ¹⁴C concentrations of 90 ± 10 pMC and ¹⁸O values of $-6.8 \pm 0.4\%$, as expected for present-day precipitation. Deeper bores discharge water with lower ¹⁴C concentrations and more negative ¹⁸O values resulting from input of much older water from depth. The deep end-member of the mixing trend is identified as paleowater (termed the "glacial" component) with ¹⁴C concentrations observed in the ¹⁸O values of $-6.8 \pm 0.4\%$, and chemical concentrations of the waters.

Identification of the deep groundwater as glacial water suggests that there may be a large body of such water onshore and offshore at deep levels. This water would have been caught by rising sea levels that greatly reduced the rate of flow in the aquifer. More generally, the influence of changing sea levels in the recent past (geologically speaking) on the disposition of groundwaters in coastal areas of New Zealand may have been far greater than we have previously realized.

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REFERENCES

- Clark ID, Fritz P. 1997. Environmental Isotopes in Hydrogeology. New York: Lewis Publishers. 328 p.
- Edmunds WM. 2001. Palaeowaters in European coastal aquifers—the goals and main conclusions of the PALEAUX project. In: Edmunds WM, Milne CJ, editors. *Palaeowaters in Coastal Europe: Evolution of Groundwater Since the Late Pleistocene*. London: Geological Society, Special Publications 189:1–16.
- Edmunds WM, Buckley DK, Darling WG, Milne CJ, Smedley PL, Williams AT. 2001. Palaeowaters in the coastal regions of southern and eastern England. In: Edmunds WM, Milne CJ, editors. *Palaeowaters in Coastal Europe: Evolution of Groundwater Since the Late Pleistocene.* London: Geological Society, Special Publications 189:71–92.
- Hendy CH, Wilson AT. 1968. Palaeoclimatic data from speleothems. *Nature* 216:48–51.
- Hulston JR, Taylor CB, Lyon GL, Stewart MK, Cox MA. 1981. Environmental isotopes in New Zealand hydrology. Part 2. Standards, measurement techniques and reporting of measurements for oxygen-18, deuterium and tritium in water. *New Zealand Journal of Science* 24:313–22.
- Lihou Joanne C. 1992. Reinterpretation of seismic reflection data from the Moutere Depression, Nelson region, South Island, New Zealand. *New Zealand Journal of Geology and Geophysics* 35:477–90.

- Loosli HH, Aeschbach-Hertig W, Barbecot F, Blaser P, Darling WG, Dever L, Edmunds WM, Kipfer R, Purtschert R, Walraevens K. 2001. Isotopic methods and hydrogeological context in the investigation of palaeowaters. In: Edmunds WM, Milne CJ, editors. *Palaeowaters in Coastal Europe: Evolution of Groundwater Since the Late Pleistocene*. London: Geological Society, Special Publications 189:193– 212.
- Rattenbury MS, Cooper RA, Johnston MR, compilers. 1998. Geology of the Nelson Area. Institute of Geological and Nuclear Sciences 1:250,000 geological map 9. (1 sheet + 67 p) Lower Hutt, New Zealand.
- Rosen MR. 2001. Hydrochemistry of New Zealand's aquifers. In: Rosen MR, White PA, editors. *Ground-waters of New Zealand*. Wellington: New Zealand Hydrological Society Inc. p 77–110.
- Shackleton NJ. 1987. Oxygen isotopes, ice volume and sea level. *Quaternary Science Reviews* 6:183–90.
- Stevens GR. 1990. Rugged landscape: the geology of central New Zealand, including Wellington, Wairarapa, Manawatu and the Marlborough Sounds. Wellington: DSIR Publishing. 286 p. (DSIR information series, ISSN 0077-9636:169).
- Stewart MK, Dicker MJI, Johnston MR. 1981. Environmental isotopes in New Zealand hydrology. Part 4. Oxygen isotope variations in subsurface waters of the

Waimea Plains, Nelson. *New Zealand Journal of Science* 24:339–48.

- Stewart MK, Morgenstern U. 2001. Age and source of groundwater from isotope tracers. In: Rosen MR, White PA, editors. *Groundwaters of New Zealand*. Wellington: New Zealand Hydrological Society, Inc. p 161–83.
- Stewart MK, Thomas JT. 2002. Moutere Valley groundwater: nature and recharge from isotopes and chemistry. *Institute of Geological and Nuclear Sciences Science Report 2002/22*. 28 p.
- Taylor CB. 1994. The relationship between electrolytic deuterium and tritium separation factors, and attainment of improved accuracy in radiometric low-level tritium measurement. *Journal of Applied Radiation and Isotopes* 45(6):683–92.
- Taylor CB, Evans CM. 2000. Isotopic indicators for groundwater hydrology in Taranaki, New Zealand. *Journal of Hydrology (NZ)* 38(2):237–70.

- Taylor CB, Trompetter VJ, Brown LJ, Bekesi G. 2001. The Manawatu aquifers, North Island, New Zealand: clarification of hydrogeology using a multidisciplinary environmental tracer approach. *Hydrological Processes* 15:3269–86.
- Thomas JT. 1989. Hydrogeology of the Moutere Valley, Nelson, New Zealand [Unpublished MSc Eng. Geol. Thesis]. Department of Geology, University of Canterbury, New Zealand.
- Thomas JT. 1991. Moutere Valley groundwater investigations 1989/1990. Unpublished Nelson Marlborough Regional report. 81 p.
- Thomas JT. 1992. Hydrogeology of the Moutere Valley with emphasis on groundwater recharge. Unpublished Tasman District Council report. 8 p.
- Thomas JT. 2001. Groundwater resources of the Tasman Region. In: Rosen MR, White PA, editors. *Groundwa*ters of New Zealand. Wellington: New Zealand Hydrological Society, Inc. p 411–25.