# Palaeoproterozoic magnesite: lithological and isotopic evidence for playa/sabkha environments

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#### ABSTRACT

Magnesite forms a series of 1- to 15-m-thick beds within the  $\approx 2.0$  Ga (Palaeoproterozoic) Tulomozerskaya Formation, NW Fennoscandian Shield, Russia. Drillcore material together with natural exposures reveal that the 680m-thick formation is composed of a stromatolite-dolomite-'red bed' sequence formed in a complex combination of shallow-marine and non-marine, evaporitic environments. Dolomite-collapse breccia, stromatolitic and micritic dolostones and sparry allochemical dolostones are the principal rocks hosting the magnesite beds. All dolomite lithologies are marked by  $\delta^{13}$ C values from +7.1‰ to +11.6‰ (V-PDB) and  $\delta^{18}$ O ranging from 17.4‰ to 26.3‰ (V-SMOW). Magnesite occurs in different forms: finely laminated micritic; stromatolitic magnesite; and structureless micritic, crystalline and coarsely crystalline magnesite. All varieties exhibit anomalously high  $\delta^{13}$ C values ranging from +9.0% to +11.6% and  $\delta^{18}$ O values of 20.0-25.7%. Laminated and structureless micritic magnesite forms as a secondary phase replacing dolomite during early diagenesis, and replaced dolomite before the major phase of burial. Crystalline and coarsely crystalline magnesite replacing micritic magnesite formed late in the diagenetic/metamorphic history. Magnesite apparently precipitated from sea water-derived brine, diluted by meteoric fluids. Magnesitization was accomplished under evaporitic conditions (sabkha to playa lake environment) proposed to be similar to the Coorong or Lake Walyungup coastal playa magnesite. Magnesite and host dolostones formed in evaporative and partly restricted environments; consequently, extremely high  $\delta^{13}$ C values reflect a combined contribution from both global and local carbon reservoirs. A <sup>13</sup>C-rich global carbon reservoir ( $\delta^{13}$ C at around +5%) is related to the perturbation of the carbon cycle at 2.0 Ga, whereas the local enhancement in  ${}^{13}C$  (up to +12%) is associated with evaporative and restricted environments with high bioproductivity.

**Keywords** Carbon, dolomite, isotopes, magnesite, oxygen, Palaeoproterozoic, playa, red beds, sabkha, stromatolite.

# **INTRODUCTION**

Despite the fact that magnesite is a rare mineral in sedimentary rocks, it forms large-scale deposits,

of which only two major types are exploited at present: ultramafic-hosted deposits of cryptocrystalline magnesite ('Kraubath type') and deposits of sparry magnesite within ancient marine platform carbonates ('Veitsch type'). Magnesite in lacustrine sediments in the vicinity of ultramafic rocks ('Bela Stena type') and metamorphosed ultramafic rocks with elevated magnesite content ('Greiner type') are of minor economic interest. In recent environments, magnesite occurs in coastal lakes and sabkhas and in continental playa lakes (Pohl, 1989).

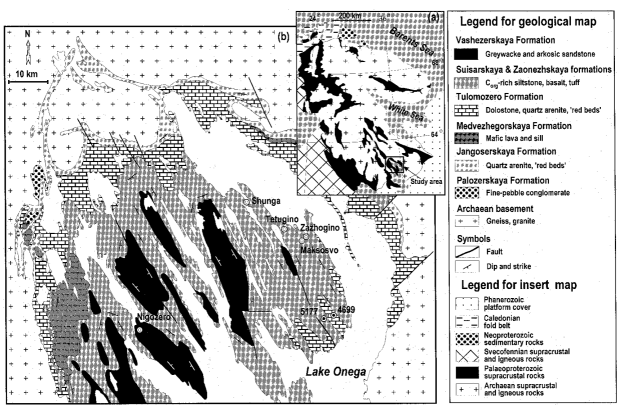
Sedimentary-hosted magnesite deposits, being an exclusive feature of Neoproterozoic–Palaeozoic marine shelf sediments, may be considered, along with abundant associated dolomite, to be an important component of Earth's evolutionary history. Archaean (Schidlowski *et al.*, 1975) and Palaeoproterozoic (Aharon, 1988) sedimentaryhosted magnesites have been described from very few localities. Magnesite mineralization from Palaeoproterozoic sedimentary environments of the Fennoscandian Shield has not yet been reported, although the possible presence of magnesite in Palaeoproterozoic dolostone sequences has long been recognized (e.g. Aksenov *et al.*, 1975).

In this article, Palaeoproterozoic sedimentary magnesite mineralization from the  $\approx 2$  billionyear-old Tulomozerskaya Formation (TF) of the NW Fennoscandian Shield (Fig. 1) is reported. The magnesite mineralization occurs within the TF carbonates, which are marked by an extreme enrichment in <sup>13</sup>C reaching +18% (Yudovich *et al.*, 1991). The TF carbonate sequence is part of the 2·3–2·06 Ga Palaeoproterozoic positive carbon isotope excursion (e.g. Baker & Fallick, 1989; Karhu, 1993) and is marked by the greatest enrichment in <sup>13</sup>C known from the Precambrian (Melezhik *et al.*, 1999).

This paper addresses the following: (i) the nature of the magnesite mineralization; (ii) depositional environments of the host carbonate sequence; and (iii) the possible significance of magnesite for understanding the extreme <sup>13</sup>C-enrichment of the host dolostones.

# **GEOLOGICAL BACKGROUND**

The local geology, stratigraphy and lithology were described in detail by Sokolov (1987). The TF is one of seven formations in the Palaeoproterozoic succession of the N. Onega Lake area. The Palaeoproterozoic succession starts with basal polymict conglomerates (Palozerskaya Formation; Fig. 1) unconformably overlying the Archaean substratum. The basal conglomerates are con-



**Fig. 1.** (a) Geographical and geological location of the study area (marked by a square). (b) Geological map (the same area marked in the small box) of the northern Onega Lake area (simplified from Akhmedov *et al.*, 1993).

formably overlain by the 50- to 120-m-thick Jangozersakaya Formation, which consists of red terrigenous rocks characterized by cross-bedding and desiccation cracks. These rocks are conformably overlain by the 70-m-thick Medvezhegorskaya Formation, which consists of mafic lava with subordinate pale grey and red, cross-bedded quartz arenites and fine-pebble conglomerates. The beds are conformably overlain by the TF, which is a 680-m-thick unit composed of stromatolitic dolostones, red quartz arenites and siltstones. This formation is, in turn, unconformably overlain by organic carbon-rich siltstones and mudstones with subordinate dolostones of the 1500-m-thick Zaonezhskaya Formation. The latter is followed by the Suisarskava Formation, a 400-m-thick succession of basalts intercalated with numerous gabbro sills. A gabbro sill from the upper part of the Suisarskaya Formation has a Sm-Nd mineral isochron age of 1980 ± 27 Ma (Pukhtel' et al., 1992). The Palaeoproterozoic succession ends with the 190-m-thick Vashezerskaya Formation comprising greywacke and arkosic sandstones.

The entire sequence was deformed and underwent greenschist facies metamorphism during the 1.8 Ga Svekofennian orogeny. The paragenesis chlorite–actinolite–epidote reflects a temperature of 300–350 °C (Sokolov, 1987).

# LITHOSTRATIGRAPHIC POSITION OF MAGNESITE BEDS

The magnesites cannot be distinguished from the host dolostones in hand samples and have been found as a result of systematic analysis. Five beds of magnesite and magnesite-bearing dolostones have been detected (Fig. 2). Four layers are in drillhole 5177 at depths of 553 $\cdot$ 5, 568 $\cdot$ 5, 598 $\cdot$ 7 and 799 $\cdot$ 0 m. In drillhole 4699, magnesite-bearing dolostones occur at depths of 537 $\cdot$ 5 m and 758 $\cdot$ 0 m (Fig. 2). Most of the beds are apparently <2 m thick, whereas the major magnesite bed (5177-553 $\cdot$ 5) is approximately 15 m thick. The lateral extent of magnesite-bearing beds is unknown. The thickest magnesite bed is sandwiched between two beds of dissolution-collapse breccia (Fig. 2).

# PETROGRAPHIC DESCRIPTION OF MAGNESITE

The magnesite and magnesite-rich dolostone are white, grey or yellow. The magnesite beds are

mainly composed of structureless micritic (5177-568·5, Fig. 3), crystalline (5177-553·5, Fig. 4) and coarsely crystalline varieties. Finely laminated micritic, stromatolitic magnesite is rare (5177-598·7).

The structureless micritic magnesite consists of micritic magnesite  $(5-10 \mu m)$  with remnants of dolomite. Rare idiomorphic crystals (0.1–0.3 mm) of magnesite and dolomite developed in the micritic aggregate (Fig. 3) indicate recrystallization related to neomorphic/metamorphic processes. Micritic magnesite is also found in sparry allochemical dolostones with syntaxial dolomite cement. Here, small magnesite crystals are present within the dolomitic intraclasts and are absent in the syntaxial dolomite cement and crystalline dolomite matrix. If clear syntaxial overgrowths formed mainly in a burial environment (e.g. Tucker & Wright, 1990), then the magnesite crystals probably replaced dolomite before most of the burial carbonate cements formed.

Micritic magnesite replaced dolomite pseudomorphically, producing very finely intergrown magnesite-dolomite aggregate (Fig. 4). From the micritic dolomite-magnesite relationship, it is obvious that magnesite could only have originated during early diagenesis if dolomite is a primary precipitate. Alternatively, magnesites may have formed somewhat later if dolomite is an early diagenetic replacement of a calcite precursor.

The structureless crystalline magnesite consists of 0.1-0.5 mm idiomorphic or subidiomorphic magnesite crystals apparently replacing either micritic aggregate of 15–10 µm magnesite crystals (Fig. 4) and subordinate dolomite rhombs or silty dolomitic micrite. The coarsely crystalline magnesite has a larger grain size (0.5-2.0 mm) but exhibits similar relationships to the earlier phases of micritic dolomite and magnesite.

Micritic stromatolitic magnesite has undergone recrystallization. When fine lamination is preserved, the thin, pale grey laminae are predominantly composed of micrite, whereas thicker, light laminae consist of microsparite.

# LITHOFACIES AND DEPOSITIONAL ENVIRONMENTS

Ten lithofacies are recognized in the TF sequence (Table 1, Fig. 2). These lithofacies and their depositional environments have been described by Melezhik *et al.* (2000), who demonstrated that the TF was deposited in a variety of environments. Terrigenous 'red beds' present throughout

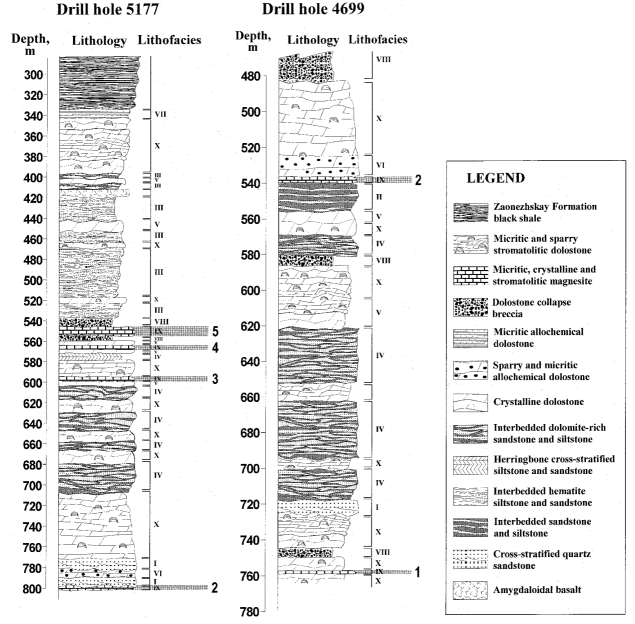
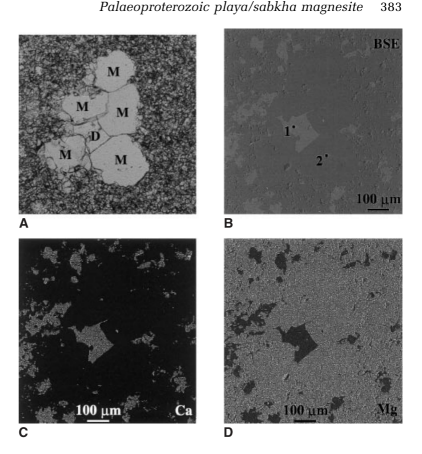


Fig. 2. Lithofacies and position of magnesite and magnesite-bearing dolostone beds in the TF intersected by drillholes 5177 and 4699. Magnesite beds are marked by grey bars.

the sequence formed in three main depositional settings: (i) a braided fluvial system over a lowenergy, river-dominated coastal plain (lithofacies I); (ii) a low-energy, barred, evaporitic lagoon (lithofacies II and IV–VII); and (iii) a non-marine, playa lake (lithofacies III and VIII). Biostromal and biohermal columnar stromatolitic dolostones (lithofacies X) are abundant and formed in shallow-water, low-energy, intertidal zones, barred evaporitic lagoons and peritidal evaporitic environments. The red, flat-laminated stromatolites formed in evaporative ephemeral ponds, coastal sabkhas and playa lakes. The presence of tepees, mudcracks, halite casts, pseudomorphs after calcium sulphate and abundant 'red beds' in the sequence suggests that terrestrial environments dominated over aqueous, with partial or total decoupling between the stromatolite-dominated depositional systems and nearby sea.

The magnesite has been designated as lithofacies IX (Melezhik *et al.*, 2000).

Further details will only be given for those lithofacies that are closely associated with the five magnesite beds documented.



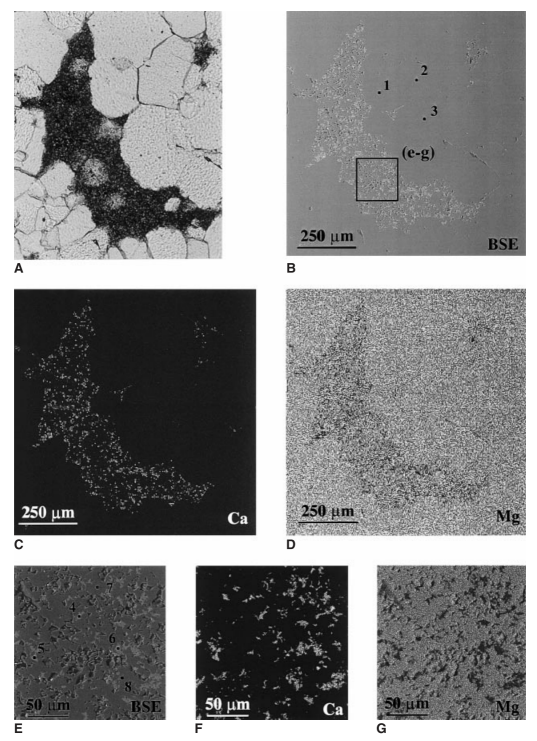
**Fig. 3.** Photomicrographs showing large neomorphic crystals of magnesite (marked by 'M') and dolomite (marked by 'D') and micritic magnesite replacing micritic dolomite. (A) Plane-polarized light; (B) back-scattered electron image; (C) Ca X-ray map; (D) Mg X-ray map. Drillhole 5177, depth 568.5 m. 1 and 2 in (B) are positions of electron microprobe analyses shown in Table 3.

Bed 5 (Fig. 2) is the major magnesite occurrence. It is sandwiched between the dissolution-collapse breccia of lithofacies VIII. The magnesite-collapse breccia assemblage is overlain by lithofacies III, which consists of intercalated brown, mudcracked, haematite-rich siltstones and pink, platy, cross-stratified, dolomite-cemented quartz arenites. Fine laminae in lithofacies III typically form low-relief hummocks and swells with amplitudes of less than 1 cm and wavelengths of 1-3 cm. A 1-to 2-m-thick bed of clastic haematite ore and halite cube casts in siltstones have also been reported (Akhmedov et al., 1993). Lithofacies III is similar to the terrestrial 'red bed'-dolostonehalite association in the Bitter Spring Formation in the Amadeus Basin, Australia, which formed in a series of shallow, hypersaline lakes and ponds (Southgate, 1986).

The presence of halite casts and the absence of calcium sulphate evaporites is also consistent with a non-marine origin for lithofacies III. The abundant desiccation cracks and the presence of small-wavelength wave ripples suggest deposition in a playa lake.

The dissolution-collapse breccia, which hosts the Bed 5 magnesite, appears as poorly cemented fragments of brown, pink and white dolostones and brown, finely laminated mudstones either embedded in insoluble residues or cemented by coarsely crystalline dolomite (Fig. 5A). The insoluble residues are composed of dark brown dolomite-sericite-chlorite material enriched in iron oxide. As no palaeokarst surfaces have been observed, the subsurface dissolution of evaporite minerals was probably the main process that led to the development of the collapse breccias. Sedimentological data match a playa lake or sabkha environment (Melezhik *et al.*, 2000). The close spatial relationship of Bed 5 to lithofacies III and VIII indicates that it formed either in a playa lake or in a ponded tidal flat setting under evaporitic conditions.

Bed 3 rests on lithofacies V and is overlain by lithofacies X. Bed 4 is entirely associated with lithofacies V, which consists of variegated, structureless or indistinctly parallel-laminated dolostones, marls and mudstones. The rocks are sporadically marked by desiccation cracks and by dolomite pseudomorphs after displacive, isolated, small crystals of gypsum (Akhmedov *et al.*, 1993). Some of these pseudomorphs have 'swallow tail' twin morphology (Fig. 5B). The desiccation cracks are filled with quartz sand, are several decimetres wide and penetrate 2–3 cm.



**Fig. 4.** Photomicrographs showing large crystals of magnesite replacing aggregate of micritic magnesite and dolomite. (A) Plane-polarized light; (B) back-scattered electron image; (C) Ca X-ray map; (D) Mg X-ray map. (E–G) Magnified views of intergrown magnesite-dolomite mixture. (E) Back-scattered electron image; (F) Ca X-ray map; (G) Mg X-ray map. Drillhole 5177, depth 553.5 m. 1–3 in (B) and 4–8 in (E) are positions of electron microprobe analyses shown in Table 3.

Syndepositional deformation is expressed as tepee structures (Akhmedov *et al.*, 1993). The lack of post-depositional compaction of the desiccation cracks suggests that the lithification occurred before burial (Melezhik *et al.*, 2000). This is consistent with the depositional environment,

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	Suggested palaeoenviron- Specific structural feature mental interpretation	1 Tepee, tepee-related See text breccias, desiccation cracks	Ι	rite, – Subsurface dissolution of halite/sulphate in playa, sabkha environment	Allochems are exclusively I dolomitic oolites	Gypsum pseudomorphed Lo by cauliflower-like quartz aggregates	Ď	–	Mud cracks, low-relief Playa or ponded tidal flat hummocks, small under evaporitic wave ripples, halite conditions casts	Ι	te Channels, tabular sets of Braided fluvial system on a cross-stratification, carbonate coastal plain asymmetric ripples
	Cement	Not preserved	Not preserved	Sericite, chlorite, haematite	Not preserved	Syntaxial dolomite spar	Not preserved	Dolomite	Dolomite	Dolomite, quartz	Quartz, sericite
interpretation.	Bedding and lamination	Laminated	Massive, laminated	I	Thinly laminated	Structureless, crudely stratified	Structureless, indistinct parallel-laminated	Herringbone cross- bedded, flaser- bedded	Platy, cross-stratified	Thin-bedded or lenticular-bedded	Ripple-marked, cross-stratified or structureless
	Rock colour	Red, beige, white	White, pale yellow	Brown, red	Red	Red	Beige, pink	Beige, pale pink	Brown, red	Grey	Grey, red
lable 1. Lithofacies and their palaeoenvironmenta	Rock assemblage	Micritic and sparry stromatolitic dolostones	Micritic, crystalline and stromatolitic magnesite	Dolomite-collapse breccias	Micritic allochemical dolostones	Sparry and micritic allochemical dolostones	Crystalline dolostones	Dolomite-rich sandstones and siltstones	Haematite siltstones and sandstones	Sandstones and siltstones	Quartz arenites
I able 1	Litho- facies	X	IX	ΠΙΛ	ΠΛ	Ν	Ν	IV	Ш	п	Ι

**Table 1.** Lithofacies and their palaeoenvironmental interpretation.

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Fig. 5. (A) Dissolution-collapse breccia consisting of fragments of brown, pink, massive and finely laminated mudstones cemented by coarsely crystalline dolomite. Lithofacies VIII, drillhole 5177, depth 542.0 m. Core diameter is 42 mm. (B) Dolomite-pseudomorphed crystals of gypsum exhibiting 'swallow tail' twin morphology. Lithofacies V, drillhole 5177, depth 570.0 m. Core diameter is 42 mm.

which was affected by long-term subaerial exposure, allowing the carbonate sediments to become lithified. The observed 'swallow tail' twin morphology of dolomite pseudomorphed gypsum crystals closely resembles those reported from ancient rocks elsewhere (Rubin & Friedman, 1977; Spencer & Lowenstein, 1990). The lithofacies V gypsum appears to have precipitated penecontemporaneously with the dolomite from shallow, near-surface brines (Melezhik et al., 2000). Because intrasediment gypsum growth provides unequivocal evidence of post-depositional crystallization in an evaporitic environment (Demicco & Hardie, 1994), similar conditions are proposed for lithofacies V. Overall, the sedimentological data suggest that the micritic dolostones of lithofacies V and the associated magnesites of Beds 3 and 4 formed in a shallow-water, evaporative setting. The tepee structures suggest that the carbonate rocks were subaerially exposed for extended periods.

Bed 2 magnesite has close association with lithofacies I, II and VI. Lithofacies II does not contain reliable genetic information. However, the main diagnostic features of lithofacies I, such as the dominance of sand, lack of silty and muddy particles, unidirectional cross-stratification, generally fining-upward sequence and presence of erosional channels, are consistent with braided fluvial systems over a low-energy, river-dominated coastal, carbonate plain. Lithofacies VI is a red grainstone, which is typically structureless or crudely stratified by variations in colour and grain size. Allochems include unsorted, rounded and angular intraclasts of dolostone and sporadic haematite and siliceous oolites. Abundant vugs, voids and cauliflower-like aggregates of quartz (with both castellated margins and mammillated surfaces) with crude radial fabric are common. Given the lack of micrite in lithofacies VI grainstones, the depositional setting must have had sufficient current or wave energy to winnow away the fine matrix (Folk, 1962). However, the unsorted, mixed rounded and angular intraclasts suggest that the depositional environment has only occasionally been influenced by such currents and waves. Another diagnostic feature is abundant cauliflower-like aggregates of quartz, which are pseudomorphs after calcium sulphate nodules (Melezhik et al., 2000). The discovery of solitary anhydrite nodules, tens of millimetres to 0.25 m in diameter, in the Holocene sediments of the Persian Gulf (Curtis et al., 1963; Shearman, 1966) led to the development of 'carbonateevaporite' or 'sabkha' depositional models to explain some ancient shallow-marine carbonate deposits (Shinn, 1983; James, 1984; Hardie & Shinn, 1986). Thus, overall, the data suggest that lithofacies II and VI, and the associated magnesite Bed 2, formed in a barred, river-dominated, coastal, evaporative carbonate plain, occasionally influenced by tidal currents and waves.

Bed 1 magnesite is hosted by lithofacies X red, haematite-rich, flat-laminated stromatolites that

form relief sheets. The stromatolite sheets are very often cracked and characterized by either indistinct, clotted or ribbon fabric with polygonal prism cracks. The presence of blisters, clotted fabrics with fenestrae and abundant desiccation cracks suggests that the flat-laminated stromatolites and associated magnesite Bed 1 formed in drained depressions and ephemeral ponds on a carbonate flat (playa lake or sabkha environment).

#### SAMPLES AND ANALYTICAL METHODS

Magnesite and dolomite samples were obtained from drillcores. The drillholes 5177 (35°25'00"E, 62°14'29"N) and 4699 (35°28'00"E, 62°14'30"N) were made by the Karelian Geological Expedition. Both drillholes are 800 m deep; they partly overlap and intersect the entire thickness (680 m) of the TF.

Whole-rock oxygen and carbon isotope analyses were carried out at the Scottish Universities Environmental Research Centre using the phosphoric acid method described by McCrea (1950) and modified by Rosenbaum & Sheppard (1986) for operation at 100 °C. Carbon and oxygen isotope ratios were measured on a VG SIRA 10 mass spectrometer. Calibration to international reference material was through NBS 19, and precision  $(1 \sigma)$  for both isotope ratios is better than  $\pm 0.2\%$ . Oxygen isotope data were corrected using the fractionation factor 1.00913 for dolomites and 1.00933 for magnesite recommended by Rosenbaum & Sheppard (1986). The  $\delta^{13}$ C data are reported in per mil (%) relative to V-PDB and the  $\delta^{18}$ O data in % relative to V-SMOW.

Because the magnesite and dolomite are commonly intergrown, the chemistry (XRF) and stable isotopic composition of the samples were obtained by whole-rock analysis, with additional electron microprobe (EMP) measurements to detect small-scale chemical variation. A sequential acid reaction was used in an attempt to resolve isotopic composition of different mineralogical components in finely intergrown mixtures of magnesite and dolomite. The approach was based on the procedures recommended by Al-Aasm et al. (1990). A three-step sequential dissolution with phosphoric acid was used on 10 mg aliquots: (i) 2 h at 25 °C to react calcite; (ii) 24 h at 50 °C to react dolomite; (iii) 2 days at 100 °C to react magnesites.

The major and trace elements were analysed by X-ray fluorescence spectrometry at the Geological Survey of Norway using a Philips PW 1480 X-ray spectrometer. The accuracy  $(1 \sigma)$  is typically better than 2% of the oxide present (SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, MgO, CaO), even at the level of 0.05 wt%, and the precision is almost invariably higher than the accuracy. The analytical uncertainties  $(1 \sigma)$  for Sr, MnO and Fe<sub>2</sub>O<sub>3</sub> are better than  $\pm 5.5$  p.p.m.,  $\pm 0.003\%$  and  $\pm 0.01\%$  respectively.

Back-scattered electron imaging, EMP measurements and cathodoluminescence of carbonate minerals were carried out at the Institute of the Continental Shelf in Trondheim. A Jeol 733 SEM Microprobe (Noran Instruments) with a silicon/ lithium detector and Norwar window type were used. The operating conditions were as follows: take-off angle 40°, acceleration voltage 15 kV, beam current 15 nA, beam diameter 1 µm, working distance 11 mm. A Proza-type matrix correction (fourth generation) was used. The detection limit for Ca, Mg, Fe and Mn is 0.1 wt% (energy dispersive spectrometer). The following calibration standard references were used: calcite for Ca, dolomite for Mg, magnetite for Fe, bustamite for Mn.

# **GEOCHEMICAL RESULTS**

Fifteen whole-rock samples of dolostone and four samples of magnesite from drillholes 5177 and 4699 were analysed for oxygen and carbon isotopes as well as for major and trace elements (Table 2). Results of electron microprobe analysis carried out on the selected samples are presented in Table 3. Additionally, a sequential acid reaction with subsequent oxygen and carbon isotope analyses was used for 28 composite dolomite– magnesite samples (Table 4).

# Dolostone

Mg/Ca ratios of the dolostone dolomite, based on EMP measurements, range from 0.46 to 0.63 (Table 3) with an average of 0.55, which is lower than that for stoichiometric dolomite (0.62).

The spread in oxygen and carbon isotope values is from 17.4% to 26.3% (mean  $21.5 \pm 1.7\%$ ) and from +7.1% to +11.6% (mean  $+9.6 \pm 1.3\%$ ) respectively. Cross-plots reveal no statistically significant covariation between  $\delta^{13}$ C and  $\delta^{18}$ O (Fig. 6). There is no relationship between the Mg/Ca ratio and C/O isotopes, nor between Mn, Mn/Sr and C/O isotopic values. Sr and  $\delta^{13}$ C and Sr and  $\delta^{18}$ O show positive covariation, r = 0.70 and r = 0.48 respectively.

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Table 2. W.	Table 2. Whole-rock elemental and isotopic composition of dolomite and associated magnesite from the TF	mposit	ion of do	lomite a	nd asso	ciated n	nagnesit	e from t	he IF.						
Depth (m/Sample)	Depth (m/Sample) Lithology	SiO <sub>2</sub> (%)	Al <sub>2</sub> O <sub>3</sub> (%)	${\rm Fe_2O_3}$ (%)	MgO (%)	CaO (%)	Na <sub>2</sub> O (%)	MnO (%)	F (%)	Sr (p.p.m.)	δ <sup>13</sup> C (‰)	$\delta^{18}$ O (%)	Mg/ Ca	Ca/ Sr	Mn/ Sr
Drillhole 51	Drillhole 5177, dolostones 542.0 Discolution-collarse hracria	1. 1		0.02	91.0	31.8	0.14	0.9E	0.43	910 	0.0	18.6	0.58	1030	8.70
542.5	Dissolution-collapse breccia	- 7 7.8		0.02 0.02	21·5	30-9	0.14	0.17	0.43	213 185	0.0 0	19.6	0.59	1186	7.08
554.5	Dissolution-collapse breccia	I	Ι	0.02	22.8	32.4	0.13	0.08	0.44	169	7.9	21.8	0.59	1361	3.64
555.5	<b>Crystalline</b> dolostone	1.5	Ι	0.02	22.5	31.4	0.13	0.07	0.45	158	$8 \cdot 1$	21.8	0.61	1410	3.41
556.0	<b>Crystalline</b> dolostone	0.2	I	0.02	22.4	32.2	0.13	0.07	0.46	166	8.0	22.2	0.59	1375	3.25
560.5	<b>Crystalline</b> dolostone	7.5	I	0.02	19.3	28.3	0.14	0·07	0.40	118	9.3	21.7	0.58	1702	4.57
563.0	Stromatolitic dolostone	13.2	I	0.02	17.6	25.2	0.13	0.01	0.34	83	9.3	17.5	0.59	2152	0.93
584.5	Sparry allochemical dolostone	34.1	0.25	0.14	14.1	20.5	0.13	60.0	0.25	82	7.9	$21 \cdot 1$	0.58	1778	8.45
593.0	Stromatolitic dolostone	7.6	I	0.03	19.6	28.3	0.14	0.05	0.38	181	8·2	21.7	0.59	1109	2.13
599.0	Crystalline dolostone	3.9	0.45	0.24	22.0	29.5	0.13	0.06	0.43	193	8.3	22.5	0.63	1085	2.39
788·0	<b>Crystalline</b> dolostone	8.9	0.61	0.24	19.0	27.4	0.16	I	0.57	67	10.6	22.8	0.68	2904	I
796.5	<b>Crystalline</b> dolostone	8.7	I	0.06	19.7	27-4	0.13	0.05	0.37	318	10.7	22.0	0.59	612	1.21
Drillhole 51	Drillhole 5177, magnesite														
553.5	<b>Crystalline magnesite</b>	14.0	I	I	26.1	0·8	I	0.17	I	5	0.6	23.1	26.5	1179	261.80
568.5	Micritic magnesite	I	I	I	30.6	3.5	Ι	0.19	Ι	11	9.4	24.3	7.37	2266	133.00
598.7	Stromatolitic magnesite	8·2	1.60	0.35	26.8	9.1	I	0.05	I	62	9.6	21.4	2.50	1039	6.21
799.0	Coarsely crystalline magnesite	2.3	1.20	0.43	31.5	8.6	I	0.06	I	71	11.3	20-9	2.63	860	I
Drillhole 46	Drillhole 4699, dolostones 523.5 Screme allochamical doloctono	15.0	01.0	0.0	17.1	0.10	К Г.О	10.0	0.00	и И	0.6	9.10	0.5.0	2105	6.60
524-1 524-1	Sparry allochemical doloctone	0 01 8.2		00.0	10.4	0 47 0	115 15	10.0	0.43	50	0 0 0	0 T 7	0000 85.0	7681	0 0 6-16
527.5	Sparry allochemical dolostone	4 5 5	I	0.06	21-4 21-4	29.6	0-13 0-13	0.04	0.55	491	10.8	23.0	0.61	429	0.63
530.0	Sparry allochemical dolostone	1.5	I	0.04	22.0	31.4	0.13	0.04	0.53	449	10.9	23.3	0.59	496	0-69
535.5	Sparry allochemical dolostone	14.7	I	0.02	17.5	24.6	0.12	0.04	0.34	396	11.1	22.0	0.60	440	0·78
Drillhole 46 537·5	Drillhole 4699, magnesite-bearing dolostone 537-5 Crystalline magnesite	1.5	I	0.02	32.8	10.6	I	0.05	I	151	11.2	21.0	2.62	497	2.55
–, Below de	-, Below detection limit. Detection limit is <0.1 for		$SiO_2$ , <5 for Sr and <0.01 for other elements.	r Sr and	<0·01 f	or other	elemen	ts.							

**Table 2.** Whole-rock elemental and isotopic composition of dolomite and associated magnesite from the TF.

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	4699-506·0	C					5177-553·5	3.5						
	Dolomite allochem	allochem		Cement	Cement, syntaxial dolomite	dolomite	Crystalli	Crystalline magnesite	site		Micritic	Micritic dolomite		
Component			Average			Average	1*	2*	3★	Average	4*	5*	€★	Average
MgO*	37.93	38·22	38·08	36.59	37.46	37-03	97-96	97.17	98.67	97-93	36.10	35.73	34.25	35.36
FeO*	0.34	0.00	0.17	0.19	00.0	0.10	00.0	0.57	0.00	0.19	0.14	0.19	0.24	0.19
CaO*	60.73	61.55	61.14	62.48	61.72	62.10	0.20	0.43	0.22	0.28	60·85	62.55	63·09	62.16
$MnO^*$	0.00	0.00	0.00	0.32	0.31	0.32	0.73	0.83	0.72	0.76	0.69	0.91	0.80	0.80
Sum	00.66	99.77		99-58	99-49		98·89	00.66	99.61		97.78	99·38	98·38	
$MgCO_3$ †	42.30	42.28	42.29	40.68	41.62	41.15	99·26	98-54	99.25	99-02	40.88	39-88	38·69	39.81
$FeCO_3^{\dagger}$	0.29		0.15	0.16	0.00	0.08	00.0	0.44	0.00	0.15	0.12	0.16	0.21	0.16
$CaCO_3$	57.41	57.72	57.56	58.88	58.12	58.50	0.17	0.37	0.19	0.24	58.40	59.18	60.41	59.33
$MnCO_3$	00.0	00.0	00.0	0.27	0.27	0.27	0.57	0.65	0.56	0.59	0.60	0.78	0.70	0.69
Sum	100.00	100.00		100.00	100.00		100.00	100.00	100.00		100.00	100.00	100.00	
Mg/Ca	0.63	0.63	0.63	0.59	0.61	0.60	493·20	227.60	451.60	340	0.60	0.58	0.55	0.57
	5177-553·5	.5		2	5177-568·5		5177.	5177-584·5			5177-796-5	796-5		
	Micritic r	Micritic magnesite			Dolomite	Crystalline magnesite	Zone	d enhedra	Zoned enhedral dolomite		Micrit	Micritic magnesite	ite	
		0				0						ο		
Component	7★	8★	Average		1★	2★	Over	Overgrowth	Edge	Core			ţ	Average
MgO*	66-96	6 <i>2</i> ·79	97-39		37-57	98.82	32-32	2	35.73	33.97	98-43		98·14	98·29
FeO*	0.16	00.0	0.08		00·0	0.00	00.0	0	0.00	0.00	0.36		0.00	0.18
CaO*	0-77	0.45	0.61		61.60	0.15	66.59	6	62.36	63·94	0.92		0.91	0-92 2 2 2 2
MnU*	0.53	0.02	0.28		0.09	0.39	0.00	0,	0.94	1.12	0.04		0.41	0.23
Sum	98·45	98·26	01.00		99-26 11 21	99-36 50 77	98-91	<u> </u>	99-03 10-01	99-03	99.75 20.21		99-46	10.00
MgCU <sub>3</sub> †	98-79 010	09.66	61.66	4.	41·81 0.00	76-99 0.00	36.41	1.	40.01	38.16	19-99		98-90 6 60	98-91 0.1.1
Fecu <sub>3</sub> 7	0.12	00.0	90-0		0.00	0.00	0.00	0	0.00	0.00	0.28		0.00	0·14 2 - 2
CaCU <sub>3</sub> † MnCO +	0-66	0:39	0.53		58·11 0.08	0.13	63·59 0·00	6	59·18 0.81	60.87 0.07	0.78		0.78 0.32	0:78 0:17
	00.001	100.00	77 0		0.001	00.001	00.001		10.00	/6.0 / 00.001	00.001	1	7C.	/T.O
Sum Mg/Ca	126-90	218·80	173	ī	0.61	100.00 663.40	0.49	6	0.58 0	0.54 0.54	107·70			108
2														

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Core, 1 Component MgO* 30.87 FeO* 0.04 CaO* 66.95											
ant	Core. micritic dolomite	lomite	Recrystall Mantle, n	allized coated dolomite grain , micritic dolomite	d dolomite omite	grain	Rim. mic	tim. micritic dolomite	nite	Cement, syntaxial dolomite	Matrix, dolomite spar
		Δποποτο	×			Avenues			Avenage		7
		Average				Average			Average		
	7 31·92	31.40	35.87	35.14	36.11	35.71	32.70	31.80	32.25	38.65	38.49
		0.10	0.37	0.00	00.0	0.12	0.00	00.0	00.00	0.00	0.22
		66.94	62.40	62.36	63.10	62.62	66.07	66.80	66.44	60.09	60·92
		0.19	0.51	1.36	0.49	0.79	0.00	0.61	0.31	0.46	0.00
			99.15	98·86	0 <i>2</i> .66		98.77	99.21		99·20	99.63
MgCO <sub>3</sub> † 35·14	4 35·93	35.53	40.11	39.46	40.13	39-90	36.86	35.77	36.32	42.97	42.63
		0·08	0.32	00·0	0.00	0.11	0.00	00.0	00.00	0.00	0.19
$CaCO_3$ † 64·60		64.22	59.14	59.36	59.45	59.31	63.14	63.70	63.42	56.63	57.19
		0.16	0.44	1.18	0.42	0.68	0.00	0.53	0.26	0.39	0.00
Sum 100-00	10		100.00	100.00	100.00		100.00	100.00		100.00	100.00
Mg/Ca 0·46	§ 0·48	0.47	0.58	0.57	0.58	0.58	0.50	0.48	0.49	0.64	0.64

Table 3. Continued

'Measured values; †Calculated values;  $\star$ , Analysis points as marked on Figs 3 and 4.

#### Magnesite

The whole-rock XRF analyses indicate that the five beds of magnesite and magnesite-bearing dolostones are characterized by Mg/Ca ratios that increase upwards through the sequence from 2.6 to 26.5 (Table 2). This indicates that the concentration of magnesite increases upwards through the sequence with the culmination in upper magnesite Bed 5. The magnesites have lower Na and F contents than the dolomites (below detection limits; Table 2). They are also depleted in Sr (<5-151 p.p.m. averaging 57 p.p.m.) compared with the dolomites. However, in the case of magnesite, Sr having a larger ionic radius should substitute only for Ca (similar to dolomite; Kretz, 1982; Reeder, 1983). If the Sr concentrations are normalized to Ca, there is no difference between magnesites and host dolostones.

EMP measurements suggest that the Mg/Ca ratios of the magnesite correlate with the petrography of the rocks. Large magnesite crystals (neomorphic/metamorphic magnesite) have high Mg/Ca ratios (230–663), whereas the Mg/Ca ratios of micritic magnesite drop to 108-172, which corresponds to 0.78-0.53 CaCO<sub>3</sub> wt% (Table 3). The overall  $CaCO_3$  content of the magnesite is low, in the range 0.13-0.78 wt% (Table 3). The micritic magnesite is marked by a relatively enhanced  $CaCO_3$  content (0.53 wt% on average) compared with the crystalline and coarsely crystalline neomorphic magnesite (0.24 wt% on average). The crystalline and coarsely crystalline magnesite is relatively enriched in Mn compared with micritic magnesite.

 $\delta 13C$  and  $\delta^{18O}$  values range from +9.0% to +11.6‰ (mean +10.1 ± 1.0‰) and from 20.0‰ to 25.7‰ (mean 22.2 ± 1.7‰) respectively. Overall, the magnesites and their host dolostones are rather similar in carbon isotope ratios (Fig. 6, Table 4). However, the Bed 4 magnesite is characterized by slightly elevated  $\delta^{18}O$  values (25.0‰ on average) compared with the average host dolostone (21.3‰, Table 2) and intergrown dolomite (21.6‰, Table 4).

# EVALUATION OF DIAGENESIS AND METAMORPHISM

There is growing evidence that dolomite in the Precambrian precipitated either directly from sea water (Tucker, 1982) or by dolomitization during early diagenesis caused by waters isotopically comparable with sea water (e.g. Veizer & Hoefs,

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<b>Table 4.</b> Carbon and oxygen isotopeanalyses of composite dolomite-		Yield (%)	$\delta^{13}$ C	$\delta^{18}$ O	Yield (%)	$\delta^{13}$ C	$\delta^{18}$ O
magnesite samples using selective acid extraction.	Depth (m/sample)	Dolomite			Magnesite		
	Magnesite Bed 2						
	4699-537·5*					11.2	21.0
	4699-537·5	24	11.2	21.7	76	11.6	20.8
	4699-537·6	36	11.4	22.3	64	11.5	20.6
	4699-537·7	33	11.4	22.6	67	11.6	20.8
	4699-537·8	24	11.3	22.7	76	11.6	21.0
	$4699 - 537 \cdot 9$	38	11.3	22.1	62	11.5	20.8
	Magnesite Bed 5						
	5177-553·5*					9.0	23.1
	5177-553·5	0	-	-	100	9.2	$23 \cdot 2$
	5177-553·55	10	9·0	23.4	90	9.2	23.7
	5177-553·6	0	-	-	100	9.2	23.1
	5177-553·6	0	-	_	100	9·1	23.6
	5177-553·65	10	7.8	22.1	90	9·0	23.6
	5177-553.7	6	9.9	26.3	94	9.3	23.8
	5177-553·8	0	_	_	100	9.3	23.9
	5177-553.9	6	8.8	22.0	74	9·1	23.7
	Magnesite Bed 4 5177-568·5*					9·4	24.3
	5177-568·5	9	9·2	21.8	91	9.4 9.8	24.3 24.6
	5177-568·7	18	9 Z 9·7	$210 \\ 23.2$	91 82	9·8	24 0 25·7
	5177-568·8	15	97 9·3	23 2 22·0	85	9·8	257 25·4
	5177-568·9	15 10·5	9·5 9·6	22·0 22·0	89·5	9·8 9·7	23·4 24·8
		10.2	9.0	22.0	09.0	97	24.0
	Magnesite Bed 3						
	5177-598.7*					9.6	21.4
	5177-598.7	21	9.0	20.7	79	9.7	20.3
	5177-598.7	42	9.6	23.2	58	9.6	21.8
	5177-598.75	0	-	_	100	9.5	21.0
	5177-598.8	24	9.3	20.6	76	9.5	20.8
	5177-598.85	22.5	7.1	17.4	77.5	9.5	21.8
	5177-598.9	14	7.5	18.1	86	9.0	20.0
	5177-598.95	21	9.3	20.8	79	9.6	21.2
	Magnesite Bed 1						
	5177-799·0*					11.3	20.9
	5177-799.0	26	11.1	20.7	74	11.5	20.1
	5177-799.2	16	11.5	21.2	84	11.6	21.1
	5177-799.4	18	11.3	20.8	82	11.5	20.3
	5177-799.8	87	11.6	19.6	13	11.0	21.3

Yields are expressed as percentages of CO<sub>2</sub> released.

\*Whole rock, non-selective analyses of dolomite-magnesite samples.

1976; Veizer et al., 1992a,b). In the Holocene, several episodes in which dolomite formed as a direct precipitate from lake water have been reported from Lake Walyungup, Western Australia (Coshell et al., 1998), and the Coorong dolomite from South Australia has also been interpreted as a direct primary precipitate from lake water (Von der Borch, 1965; Rosen et al., 1988, 1989).

Most magnesite forms as a minor mineral during diagenesis in a hypersaline environment. However, in places, hydromagnesite occurs as a primary precipitate, i.e. Salda Gölü, south-western Turkey (Braithwaite & Zedef, 1994, 1996).

Petrographic data obtained from the TF suggest that the micritic magnesite could only have originated during early diagenesis if dolomite is a primary precipitate. The micritic magnesite replaced dolomite before the major portion of the burial carbonate cements formed. However, the crystalline and coarsely crystalline neomorphic magnesite that replaced micritic magnesite grew during late diagenesis and metamorphism.

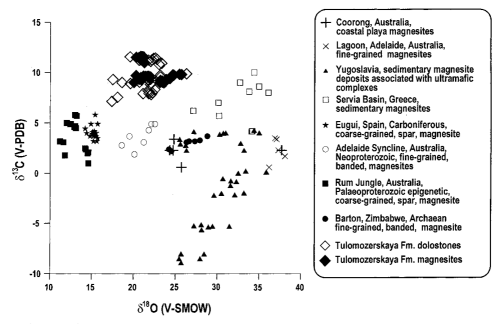


Fig. 6. Plot of  $\delta^{13}$ C vs.  $\delta^{18}$ C comparing the TF results with magnesites from other deposits. Data are from: Coorong Lagoon, South Australia (Zachmann, 1989); Lagoon, Adelaide, South Australia (Botz & von der Borch, 1984; Schroll *et al.*, 1986); magnesite deposits of Yugoslavia (Fallick *et al.*, 1991); Servia sedimentary magnesites (Kralik *et al.*, 1989); Eugui, Spain, Carboniferous, coarse-grained, spar magnesite (Kralik & Hoefs, 1978); Adelaide Syncline, Copper Claim, Australia, Neoproterozoic, fine-grained, banded magnesite (Lambert *et al.*, 1984); Rum Jungle, Northern Territory, Australia, Palaeoproterozoic, coarse-grained, spar magnesite (Aharon, 1988); Barton, Zimbabwe, Archaean, fine-grained, banded magnesite (Perry & Tan, 1972; Schidlowski *et al.*, 1975).

Several geochemical screening methods are available to assess the degree of diagenetic and metamorphic alteration. Hudson (1977) found that oxygen isotopes may be a sensitive indicator of diagenetic alteration. Diagenesis commonly decreases  $\delta^{18}$ O, and the effect of diagenesis can be revealed on a  $\delta^{13}$ C and  $\delta^{18}$ O cross-plot. Oxygen isotopes are commonly much more easily affected by exchangeable oxygen derived from either meteoric water or interstitial fluids at elevated temperatures (e.g. Fairchild et al., 1990), whereas  $\delta^{13}$ C may be buffered by the pre-existing carbonate. In general, depletion in both oxygen and carbon isotope values may be considerable during late diagenesis as well as in the course of lowgrade metamorphism accompanied by deformation (Guerrera et al., 1997).

Although a  $\delta^{13}$ C– $\delta^{18}$ O cross-plot (Fig. 6) shows no reliable correlation, the wide spread in  $\delta^{18}$ O values is evident and may reflect resetting of oxygen isotopes during later recrystallization. If the highest  $\delta^{18}$ O values of  $26 \cdot 3\%$  for dolomite and  $25 \cdot 7\%$  for magnesite are considered as the least altered, then all the lower values could have been affected by later diagenetic and metamorphic resetting. A limited spread in  $\delta^{13}$ C values suggests that carbon isotopes may have been buffered by pre-existing carbonate. It is very unlikely that fluid—rock interactions during the course of metamorphism substantially affected carbon and oxygen isotope systems, as oxygen and carbon isotope values are not significantly depleted. However, Mn enrichment in the structureless crystalline magnesite could indicate the incorporation of later diagenetic and metamorphic Mn-rich fluids.

# ORIGIN OF Mg-RICH CARBONATES FROM SEDIMENTARY ENVIRONMENTS: GENERAL

#### **Depositional environments**

Typical sedimentary magnesites are confined to: (i) ancient marine platform carbonates ('Veitsch type'; Pohl, 1989); (ii) lacustrine sediments near to or overlying ultramafic rocks (e.g. Bela Stena, Nevada; Fallick *et al.*, 1991); (iii) marine evaporates (e.g. Sebkha el Melah, east coast of Tunisia; Perthuisot, 1980); (iv) recent coastal salt flats in arid regions (e.g. sabkhas of Abu Dhabi; Bush, 1973); and (v) continental and coastal lakes (e.g. playas in Coorong Lagoon area, South Australia; Walter *et al.*, 1973; Schroll, 1989). Most magnesite forms as a minor mineral during diagenesis in a hypersaline environment. In places, hydromagnesite occurs as a primary precipitate associated with active stromatolites, i.e. Salda Gölü, south-western Turkey (Braithwaite & Zedef, 1994, 1996). The association of stromatolites and both diagenetic and primary magnesite is very common and has been reported from many recent alkaline lakes (Walter *et al.*, 1973; Last & De Deckker, 1990; Renaut, 1993; Coshell *et al.*, 1998).

# Source of Mg-rich solutions

In general, the origin of Mg-bearing solutions forming sedimentary magnesites is not well understood, although some authors suggest that it does not form as a primary phase under surface conditions (e.g. Möller, 1989). The formation of magnesite requires high Mg<sup>2+</sup>/Ca<sup>2+</sup> ratios in solution (Müller et al., 1972; Möller, 1989). It is well known that Mg<sup>2+</sup> is enriched in sea water during carbonate sedimentation. This process enhances the dolomitization of carbonate muds (Carpenter, 1980) and may proceed towards magnesitization. The influence of algae in producing high pH-values in the water may be an important factor for magnesite formation; major fossil magnesite deposits are intimately associated with biohermal stromatolitic dolomite (Misra & Valdiya, 1961; Valdiya, 1969, 1995; Raha, 1980; Shevelev et al., 1991; Joshi et al., 1993). The salinity of the diagenetic pore solutions is very important, as rising salinity enlarges the stability field of magnesite compared with that of dolomite. The Mg/Ca ratio is raised even more if sulphates crystallize (Bathurst, 1975), leading to the development of dense, Mg-enriched brines.

# Major processes controlling oxygen and carbon isotope compositions

Carbon and oxygen isotope measurements from 74 magnesite occurrences were reviewed by Kralik *et al.* (1989). These data demonstrate that cryptocrystalline–microcrystalline magnesites in ultramafic complexes are characterized by low  $\delta^{13}$ C values (-6% to -18%) and relatively high  $\delta^{18}$ O values (+22% to +29%). The negative  $\delta^{13}$ C and relatively high  $\delta^{18}$ O values indicate the formation of magnesite at low temperatures with a meteoric carbon source (Kralik *et al.*, 1989). A study of magnesite associated with ultramafic rocks of Yugoslavia by Fallick *et al.* (1991) suggested a carbon source derived from decarboxy-

lation of organic material for magnesite deposits with  $\delta^{13}C < -10^{\circ}_{\circ\circ}$ .

The fine-grained Quaternary to Recent magnesites, which occur in evaporitic sabkha and local pond environments, exhibit relatively high  $\delta^{13}$ C (+1.7‰ to +4.6‰) and  $\delta^{18}$ O values (+32‰ to +38‰). Kralik *et al.* (1989) reported that these values suggest magnesite precipitation in equilibrium with a dissolved inorganic carbon pool of ambient basinal water, with the generation of isotopically very heavy carbon by fermentation.

Ancient fine-grained magnesites show two main maxima  $\delta^{13}$ C (+2‰ to +3‰),  $\delta^{18}$ O (+25‰ to +27‰) and  $\delta^{13}$ C (-2‰ to -6‰),  $\delta^{18}$ O (+18‰ to +22‰). The <sup>13</sup>C enrichments are interpreted to result from carbon inheritance from an evaporitic carbonate precursor (e.g. Kralik & Hoefs, 1978). A mixture of evaporitic and meteoric waters has been suggested as a parent fluid for the finegrained magnesite with negative carbon and low  $\delta^{18}$ O (Kralik *et al.*, 1989).

The coarse-grained spar magnesites exhibit a wide range of  $\delta^{13}$ C (-7.5% to +4%) and  $\delta^{18}$ O (+6% to +25%). These values partly correlate with the degree of metamorphism, without evidence of magnesite formation before recrystallization (Kralik *et al.*, 1989).

Carbon and oxygen isotope compositions of sedimentary-hosted magnesites from the main types of deposits are plotted in Fig. 6 in comparison with the TF magnesite.

# ORIGIN OF Mg-RICH CARBONATES: APPLICATION TO THE TF MAGNESITE

# Mechanism of <sup>13</sup>C enrichment

The TF dolomite and magnesite exhibit extreme <sup>13</sup>C enrichment. The formation of <sup>13</sup>C-rich dolostones has been ascribed to the global 2.4-2.06 Ga positive excursion of carbonate <sup>13</sup>C/<sup>12</sup>C associated with enhanced accumulation of Corg (Baker & Fallick, 1989; Yudovich et al., 1991; Karhu, 1993; Melezhik et al., 1999). A detailed study of the TF dolostones by Melezhik et al. (1999) suggested that, although the formation of the TF <sup>13</sup>C-rich carbonates was driven by global factors, the complementary organic carbon was buried in an external basin. However, it has also been reported that TF dolostones reveal the greatest enrichment in <sup>13</sup>C ( $\delta^{13}$ C up to +18%; Yudovich *et al.*, 1991) known from this interval. Such enrichment exceeds the global value for the isotopic shift at ≈2.0 Ga (perhaps at around +5‰; Melezhik *et al.*, 1997, 1999). If the global  $\delta^{13}$ C value of +5% was caused by enhanced  $C_{org}$  burial, the further enrichment requires an extra <sup>13</sup>C-rich source(s). Development of abundant stromatolite-forming microbial communities in shallow-water basins, establishment of evaporative and partly restricted environments, high bioproductivity, enhanced uptake of <sup>12</sup>C and penecontemporaneous recycling of organic material in cyanobacterial mats with the production and consequent loss of CO<sub>2</sub> (and CH<sub>4</sub>?) have been suggested to be additional factors that may have increased  $\delta^{13}$ C from +5% up to +18% (Melezhik *et al.*, 1999).

The  $\delta^{13}$ C and  $\delta^{18}$ O values of the TF magnesite do not differ significantly from those reported for the associated TF dolostones. Thus, we assume that both magnesite and dolomite obtained their carbon from the same reservoir and a similar mechanism was involved in the carbon isotope fractionation. Therefore, the <sup>13</sup>C-rich nature of the carbon source can be assigned to both (i) the global carbon reservoir at 2.0 Ga and (ii) the local reservoir. At this stage, however, a quantitative separation between the local and global carbon reservoirs is not possible. If the global  $\delta^{13}$ C value is assumed to have been around +5% (e.g. Melezhik et al., 1997, 1999), then the  $\delta^{13}$ C of +9.0% to +11.6% can be explained as resulting from local factors, such as evaporation and restricted environments with high bioproductivity.

# Depositional environments and the source of Mg-rich solutions

As discussed previously, the depositional settings of the major magnesite units (Beds 3–5) appear to have been either sabkha to playa environments or ponds in an upper tidal flat.

Epigenetic formation of magnesite seems unlikely, as carbon and oxygen isotopes and Sr (normalized against Ca) abundances are very similar to those of the early diagenetic/sedimentary host dolomite. An external source of Mg<sup>2+</sup>rich solutions is also unlikely, as no ultramafic rocks are documented in the area. Furthermore, lacustrine magnesites formed from ultramaficderived Mg solutions exhibit a large spread in both oxygen and carbon isotope values, with the bulk of  $\delta^{13}$ C being negative (Fig. 6). Therefore, sea water is the most probable source of Mg<sup>2+</sup>-rich solutions.

The Mg<sup>2+</sup>-bearing solutions were apparently created in a hypersaline environment, as indicated by the sedimentological data. Several stages of Mg/Ca enrichment might have taken place in the basin. First, Mg/Ca may have been enriched in sea water in the course of calcium carbonate precipitation in a partly closed marine environment. Secondly, an arid climate may have led to further evaporation of sea water-derived brine as a result of gypsum, and then halite, precipitation. Progressive evaporation and salinity increases can be traced by the development of desiccation cracks and gypsum casts in the lithofacies X dolostones and then by halite casts in lithofacies III sediments, which overly the major bed of magnesite (Bed 5, Fig. 2). With rising salinity, if calcium sulphate was precipitated, the Mg/Ca ratio would have increased. The succession of these processes would have led to the development of dense, Mg<sup>2+</sup>-enriched residual brines. Depletion of magnesite in Na and F compared with dolomite may indicate deficiency in these components in magnesite-forming solutions, which is consistent with the proposition that magnesite formed from evolved brines that lost Ca and Na by earlier precipitation. Such Mg<sup>2+</sup>-rich brines would most probably have percolated downwards in the sedimentary sequence as well as laterally to the centre of the basin, causing magnesitization. Magnesite replaced dolomite, and  $\delta^{13}C$  and Sr concentration would have been buffered by the dolomite precursor, as the carbon isotope composition as well as the Ca/Sr ratios of magnesitebearing rocks are indistinguishable from those of dolomite. Petrographic data suggest that the micritic magnesite could only have originated during early diagenesis if the dolomite was a primary precipitate and replaced dolomite before the major phase of burial diagenesis. However, the crystalline and coarsely crystalline, neomorphic magnesite, which replaced micritic magnesite, grew during late diagenesis and even during metamorphism.

Relatively increased  $\delta^{18}$ O values of the magnesites compared with the  $\delta^{18}$ O average of TF dolomite are consistent with evolved evaporitic fluids. However, the original  $\delta^{18}$ O values of both dolomite and magnesite are assumed to have been partially overprinted, and the magnesite is enriched in Mn. The higher  $\delta^{18}$ O values and enhanced Mn concentration may be reconciled if a sea water-derived brine was mixed with Mnrich, <sup>18</sup>O-depleted meteoric fluids. Alternatively, the enrichment in Mn and overprinting of  $\delta^{18}$ O values could be related to a much later stage when collapse breccia formed as a result of sulphate (and halite) dissolution.

Given the limited development of magnesite in the TF, either sabkha magnesite (similar to Abu

Dhabi) or playa magnesite (similar to the Coorong Lagoon area in South Australia and Lake Walyungup in Western Australia) match available sedimentological data. The  $\delta^{18}$ O values of Coorong magnesite are similar to those of the TF, although they differ in  $\delta^{13}$ C (the magnesites studied here are richer in <sup>13</sup>C; Fig. 6). However, both magnesites and dolostones of the TF are enriched in <sup>13</sup>C; part of a more general problem related to the global  $\approx 2.0$  Ga positive excursion of carbonate carbon.

# CONCLUSIONS

1. Magnesite forms a series of 1- to 15-m-thick beds within the  $\approx 2.0$  Ga Tulomozerskaya Formation. The 680-m-thick unit is composed of a stromatolite-dolomite-'red bed' sequence formed in a complex combination of shallow-marine and non-marine, evaporitic, partly restricted environments.

2. Dolomite-collapse breccia, stromatolitic and micritic dolostones and sparry allochemical dolostones are the principal rocks hosting the magnesite beds. All dolomite lithologies have enriched  $\delta^{13}$ C values of +7.1% to +11.6% and  $\delta^{18}$ O ranging from 17.4% to 26.3%.

3. Magnesite occurs in different forms: structureless micritic, crystalline, coarsely crystalline and finely laminated micritic, stromatolitic magnesite. All varieties exhibit positive, highly anomalous  $\delta^{13}$ C values ranging from +9.0% to +11.6% and  $\delta^{18}$ O values of 20.0–25.7%.

4. Micritic magnesite originated during early diagenesis and replaced dolomite before the major phase of burial. Crystalline and coarsely crystalline, neomorphic magnesite, which replaced micritic magnesite, formed during late diagenesis/metamorphism. Magnesite apparently precipitated from sea water-derived brine, perhaps diluted by meteoric fluids. Magnesitization was accomplished under evaporitic conditions (sabkha to playa lake environment), similar to the Coorong or Lake Walyungup coastal playa magnesite.

5. Extremely high  $\delta^{13}$ C values of magnesite and dolostones probably reflect a combined contribution from both global and local carbon reservoirs. A <sup>13</sup>C-rich global carbon reservoir ( $\delta^{13}$ C at around +5‰) is related to a perturbation of the carbon cycle at 2.0 Ga, whereas the local enhancement in <sup>13</sup>C (up to +12‰) was associated with evaporative and restricted environments with high bioproductivity.

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