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# Paratethys pacing of the Messinian Salinity Crisis: Low salinity waters contributing to gypsum precipitation?



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#### ABSTRACT

During the so-called Messinian Salinity Crisis (MSC: 5.97-5.33 Myr ago), reduced exchange with the Atlantic Ocean caused the Mediterranean to develop into a "saline giant" wherein  $\sim$ 1 million km³ of evaporites (gypsum and halite) were deposited. Despite decades of research it is still poorly understood exactly how and where in the water column these evaporites formed. Gypsum formation commonly requires enhanced dry conditions (evaporation exceeding precipitation), but recent studies also suggested major freshwater inputs into the Mediterranean during MSC-gypsum formation. Here we use strontium isotope ratios of ostracods to show that low-saline water from the Paratethys Seas actually contributed to the precipitation of Mediterranean evaporites. This apparent paradox urges for an alternative mechanism underlying gypsum precipitation. We propose that Paratethys inflow would enhance stratification in the Mediterranean and result in a low-salinity surface-water layer with high Ca/Cl and SO<sub>4</sub>/Cl ratios. We show that evaporation of this surface water can become saturated in gypsum at a salinity of  $\sim$ 40, in line with salinities reported from fluid inclusions in MSC evaporites.

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#### 1. Introduction

The deposition of salts ("evaporites") on Earth is generally associated with hydrographically isolated basins in evaporative environments (Warren, 2010). By ongoing evaporation, sea water salinity increases and ultimately saturation with respect to certain minerals causes mineral deposition. The most common evaporite, gypsum (CaSO<sub>4</sub>·2H<sub>2</sub>O), forms in waters that are 4 to 5 times more concentrated in salts than average seawater:  $\sim\!130\text{-}160$  g/l (Warren, 2010).

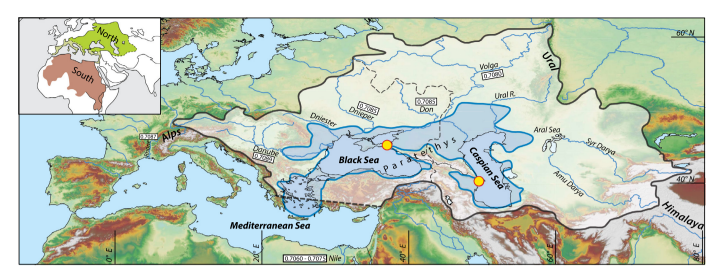
Occasionally in Earth's history, evaporite deposition took place at large scale in so-called "saline giants". Earth's most recent saline giant occurred in the Mediterranean Basin during the late Miocene (Hsü et al., 1973; Ryan, 2009). Restriction of seawater exchange between the (proto) Mediterranean Sea and the Atlantic Ocean cul-

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minated in the so-called Messinian Salinity Crisis (MSC) between 5.97-5.33 Ma (Roveri et al., 2014a; Flecker et al., 2015). The MSC is subdivided into three main stages, all characterized by specific environmental conditions (Roveri et al., 2014a). MSC Stage 1 (5.97-5.6 Myr ago) is marked by rhythmic alternations of marine marls and gypsum in the shallow basins, while the Mediterranean remained connected to the Atlantic Ocean. During MSC Stage 2 (5.6-5.55 Myr ago), the most saline conditions were reached and halite formed in the deeper basins. MSC Stage 3 (5.55-5.33 Myr ago) is characterized by large temporal salinity differences; both rhythmic gypsummarl couplets and evaporite-free fresh to brackish water deposits

Fundamental questions remain about the trigger for gypsum formation and where in the water column it formed (De Lange and Krijgsman, 2010; Lugli et al., 2010). Classically, MSC-gypsum formation is considered to result from enhanced (seasonal) drought during precession maxima (Krijgsman et al., 2001; Topper and Meijer, 2013; Topper et al., 2014; Roveri et al., 2014b). Stable



**Fig. 1.** Elevation map showing the extent of the late Miocene Black and Caspian Sea and their drainage basins. The former Paratethys Sea during the latest Miocene is outlined according to Popov et al. (2006). These authors consider the Aegean domain as an intermittent region between Mediterranean and Paratethys. Yellow dots indicate the studied sections; Zheleznyi Rog (Russia) and Adzhiveli (Azerbaijan). The main rivers are labelled with their modern <sup>87</sup>Sr/<sup>86</sup>Sr ratios. The areal distribution of the two main water sources, northern (European) and southern (African) rivers draining into the Mediterranean-Paratethys system, is presented in the inset. (For interpretation of the colours in the figure(s), the reader is referred to the web version of this article.)

**Table 1**Present-day strontium values of major Paratethyan and Mediterranean fresh water sources.

| River                          | Q<br>(km³/yr) | Sr conc.<br>(µmol/kg) | Sr flux                   | <sup>87</sup> Sr/ <sup>86</sup> Sr | Source   |
|--------------------------------|---------------|-----------------------|---------------------------|------------------------------------|--|
| Danube                         | 207           | 2.76                  | 5.71 × 10 <sup>8</sup>    | 0.7089                             | Palmer and Edmond, 1989                                |
| Dnieper                        | 43            | 2.50                  | $1.08 \times 10^{8}$      | 0.7084                             | Palmer and Edmond, 1989                                |
| Don                            | 26            | 2.50                  | $6.50 \times 10^{7}$      | 0.7084                             | Palmer and Edmond, 1989                                |
| Volga                          | 239.7         | 5.48                  | $1.31\times10^{9}$        | 0.708083                           | Clauer et al., 2000<br>Kroonenberg et al., 2011        |
| Hydrothermal vents             | ~14.6         | ~106                  | $\sim 1.55 \times 10^{9}$ | 0.70821                            | Clauer et al., 2000                                    |
| Mediterranean seawater stage 1 | _             | 87.0?                 | -                         | $\sim$ 0.7089-0.7090 $^{^{*}}$     | Roveri et al., 2014a, 2014b<br>Broecker and Peng, 1982 |
| Nile                           | -             | -                     | $1.56\times10^8$          | 0.706-0.7071                       | Brass, 1976<br>Gerstenberger et al., 1997              |
| Rhone                          | 55            | 5.94                  | $3.27 \times 10^{8}$      | 0.7087                             | Palmer and Edmond, 1989                                |
| Atlantic inflow                | 22706         | 87.0                  | $1.97\times10^{12}$       | ~0.7090*                           | Broecker and Peng, 1982<br>Bryden et al., 1994         |
| Black Sea inflow               | 300           | _                     | _                         | _                                  | Jarosz et al., 2013                                    |
| Black Sea outflow              | 400           | _                     | _                         | _                                  | Jarosz et al., 2013                                    |

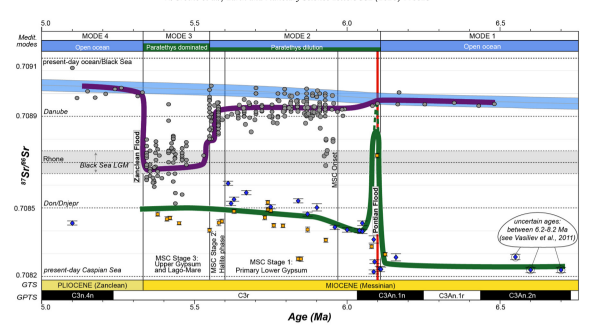
<sup>\*</sup> Messinian Salinity Crisis-values.

isotope analyses of gypsum hydration water and the salinity of fluid inclusions in MSC-gypsum, however, indicate large freshwater inputs during gypsum formation in both Stage 1 (Natalicchio et al., 2014; Evans et al., 2015) and Stage 3 (Costanzo et al., 2019). Freshwater inputs were initially attributed to Mediterranean rivers (Fig. 1 and Table 1), because the adjacent Black Sea was thought to have desiccated during the MSC (Hsü and Giovanoli, 1980; Gillet et al., 2007). Recent paleoenvironmental reconstructions, however, demonstrated that the Black and Caspian Seas remained waterfilled and had a positive hydrological budget throughout the MSC (Grothe et al., 2014; Marzocchi et al., 2016; Van Baak et al., 2017). They were unified in the large Paratethys Sea (Fig. 1), which provided a potential source for low-salinity waters to the Mediterranean (Krijgsman et al., 2010; Van Baak et al., 2015).

Episodic freshwater input into the Mediterranean is important, as it will have affected saturation state with respect to different minerals. Radiogenic strontium isotope ratios (<sup>87</sup>Sr/<sup>86</sup>Sr) are potentially a powerful tracer of Mediterranean water sources (Flecker and Ellam, 2006; Topper et al., 2014). In restricted basins, the <sup>87</sup>Sr/<sup>86</sup>Sr ratio is offset from coeval ocean water signatures and controlled by mixing with non-marine Sr sources such as hydrothermal vents and river waters (Vasiliev et al., 2010; Roveri et al., 2014b; Reghizzi et al., 2017). These regional inputs have char-

acteristic isotopic compositions and concentrations reflecting the lithology of the hinterland. During the MSC, the Mediterranean represents an archetype of a strongly restricted basin. The progressive shift from pre-MSC marine values (~0.7090) to lower values (down to 0.7085 during MSC Stage 3) (Fig. 2) is considered a further proof of the increasing deterioration of the Atlantic connection, accompanied by an increased sensitivity to continental runoff (Roveri et al., 2014b; Reghizzi et al., 2018). This provides the Mediterranean low radiogenic Sr in accordance with the Rbdepleted Mesozoic carbonates and Cenozoic basalts forming the catchment of the main peri-Mediterranean rivers (Brass, 1976; Gerstenberger et al., 1997). The role of Paratethyan waters in explaining these deviations remains uncertain as <sup>87</sup> Sr/<sup>86</sup>Sr data from the late Messinian Paratethys are lacking (Vasiliev et al., 2010).

To test whether input of Paratethyan waters into the Mediterranean potentially governed the MSC, we determined late Miocene <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the Black and Caspian waters by measuring <sup>87</sup>Sr/<sup>86</sup>Sr ratios of calcareous ostracod shells. Calcareous organisms incorporate Sr directly into their shell, thereby representing a proxy of the isotopic composition of the water they lived in (Flecker and Ellam, 2006). To reconstruct the strontium isotope composition (<sup>87</sup>Sr/<sup>86</sup>Sr) of the Paratethys Sea during the MSC, we used ostracods from two outcrop sections: Zheleznyi Rog (Black



**Fig. 2.** Late Miocene-earliest Pliocene  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  data for the Mediterranean (grey circles), Black (blue diamonds) and Caspian (orange squares) Seas. Mediterranean and oceanic (blue shaded band)  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  data are derived from various sources; see Roveri et al. (2014a) for an overview. Green and purple lines indicate our interpretation of the  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  ratio trend of the Black and Caspian Sea combined, and the Mediterranean, respectively. Samples predating  $\sim$ 6.2 Ma have large age-uncertainties because of several hiatuses in the Maeotian part of the section (Vasiliev et al., 2011). Dashed lines indicate the  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  of the Danube, Don, Dnieper, Rhone rivers, and the present-day Caspian and Black Seas.  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  ratios of the Black Sea during last glacial maximum (LGM). Messinian Salinity Stages 1-3 are labelled and Global Time Scale (GTS) and Geomagnetic Polarity Time Scale (GPTS) are shown at the bottom.

Sea) and Adzhiveli (Caspian Sea) (Vasiliev et al., 2011; Van Baak et al., 2016).

#### 2. Sections in the Paratethys

The Black and Caspian Seas were, since the Oligocene, part of the large epicontinental Paratethys Sea (Fig. 1), which covered large parts of Eurasia stretching from Germany to Tajikistan (Popov et al., 2006). Due to restricted conditions, endemic fauna developed, which form the basis for regional stratigraphic correlation. The MSC corresponds in the Black Sea to the regional Pontian Stage and part of the Kimmerian Stage (Krijgsman et al., 2010). In the Caspian region, it includes the entire Pontian Stage (Van Baak et al., 2016). The Pontian Stage is preceded by the Maeotian Stage and succeeded by the Kimmerian Stage (Black Sea) and the Productive Series (Caspian Sea).

The ~500-m-thick Zheleznyi Rog section (45°06.9082′N, 36°45.646′E), comprises ca. 5 Myr of sediments and is one of the most extensively studied Neogene successions of the Black Sea Basin (Popov et al., 2006, 2016; Krijgsman et al., 2010; Vasiliev et al., 2011; Stoica et al., 2016). The deposits of the upper Maeotian comprise grey clays and whitish diatomites, while alternations of grey marls and whitish diatomite layers characterize the Pontian. The onset of the Kimmerian Stage (ca. 5.6 Ma) is marked by an iron-rich layer of oolithic sandstone that may represent a hiatus (Krijgsman et al., 2010; Vasiliev et al., 2011). On top of this red bed, dark grey marls with interbedded limonite-rich layers mark the Kimmerian. Age constraints were provided by magnetostratigraphy coupled with biostratigraphy (ostracods, mollusks), event stratigraphy (foraminifera) and <sup>40</sup>Ar/<sup>39</sup>Ar ages (Vasiliev et al., 2011).

The Adzhiveli section (40°18.440′N, 49°4.211′E) is a 130 m thick section located in central Azerbaijan (Fig. 1). The section spans the upper Maeotian and Pontian Stages. The deposits comprise (silty) clays and marls that alternate in colour between brownish grey and dark grey (Van Baak et al., 2016). The lower part of the section includes some diatomitic beds. Marine sedi-

ments at the base of the section yield foraminifera and radiolarians that correspond to the Pontian Flooding. Based on magnetoand cyclostratigraphy coupled with biostratigraphy (ostracods) and event stratigraphy (foraminifera), this section spans roughly from  $\sim$ 6.16 Ma to <5.38 Ma (Van Baak et al., 2016).

## 3. Methods

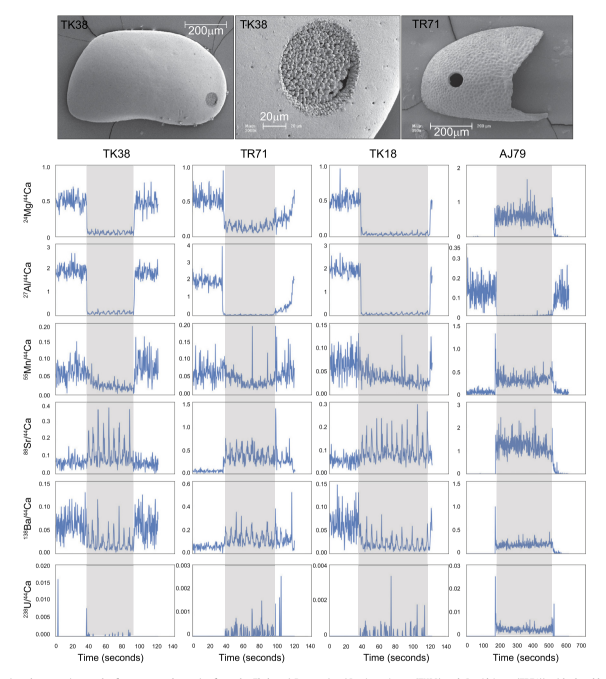
#### 3.1. Sample preparation

All  $^{87}$ Sr/ $^{86}$ Sr analyses were performed on handpicked ostracods that were separated from the bulk sediment by disaggregation in sodium carbonate solution and wet sieving to retain the >250  $\mu$ m fraction. The ostracod samples were not crushed prior to the cleaning procedure. Clays were removed by washing five times in MilliQ<sup>®</sup>, five times in methanol (96%) and by ~5-30 s cleaning in an ultrasonic bath.

#### 3.2. Diagenetic evaluation

Selected cleaned samples were evaluated for potential diagenetic alteration using trace element profiling of the shells and scanning electron microscopy (SEM; Fig. 3). Ostracod shells from Zheleznyi Rog were ablated using a Lambda Physik Excimer 193 nm laser with GeoLas 200Q optics coupled to a Micromass Platform quadrupole ICP-MS at Utrecht University. Ablation was performed in a mixed atmosphere of helium and argon with a repetition rate of 6 Hz and spot diameter of 80 µm. Calibration was performed against U.S. National Institute of Standards and Technology SRM 610 glass using the concentration data of Jochum et al. (2011) with <sup>44</sup>Ca as an internal standard. To reduce spectral interferences of the minor Ca isotopes (<sup>42</sup>Ca, <sup>43</sup>Ca, <sup>44</sup>Ca), a collision and reduction cell was used (Mason and Kraan, 2002).

Samples from Adzhiveli (Caspian Sea) were ablated in a He environment using a NWR193UC (New Wave Research) laser, containing a 193 nm short pulse ArF Excimer laser, coupled to a quadrupole ICP-MS (Thermo Scientific iCAP-Q) at the Royal NIOZ.



**Fig. 3.** Scanning electron micrograph of two ostracods species from the Zheleznyi Rog-section (*Caspiocypris atta* (TK38) and *Cyprideis* sp. (TR71)) with the ablation craters. Time resolved laser-ablation inductively coupled plasma-mass spectrometry data is shown for four representative samples. Values are normalized against <sup>44</sup>Ca to correct for variations in ablation rates. Typically, material is ablated at different rates causing a variation in counts per second. Some peaks on the outside of the shell, e.g. <sup>27</sup>Al/<sup>44</sup>Ca of TR71 or <sup>55</sup>Mn/<sup>44</sup>Ca of AJ79, indicate some clay minerals and/or coating that were not removed despite the cleaning procedure. However, the relative contribution of this outside contamination is very low. The Sr counts in the contamination peak of AJ79 account only for <0,1% of the <sup>88</sup>Sr content of the entire shell. Furthermore, all ratios show a stable signal without appreciable peaks inside the shell indicating pristine preservation. Hence, post-depositional alternation of the isotopes due to secondary processes can therefore be excluded. The non-contaminated part of the ostracods shells is marked by the grey interval.

Laser ablation was performed with an energy density of approximately 1 J/cm² and at a repetition rate of 6 Hz and an 80 µm spot diameter. Depth profiles were recorded using the Qtegra (Thermo Scientific) software package. Monitored masses included <sup>24</sup>Mg, <sup>27</sup>Al, <sup>44</sup>Ca, <sup>55</sup>Mn, <sup>88</sup>Sr, <sup>138</sup>Ba and <sup>238</sup>U (Fig. 3). Calibration was performed against U.S. National Institute of Standards and Technology SRM 610 glass ablated at a higher energy density (approximately 5 J/cm²) with <sup>44</sup>Ca as an internal standard using the values of Jochum et al. (2011) and assuming natural isotopic abundances. In addition, an in-house matrix-matched standard, based on an Iceland spar was ablated at lower energy density (1 J/cm²) with

known Mg and Sr concentrations (Raitzsch et al., 2010) to check for the effect of changing energy densities. The constant elemental ratios in the ablation profiles with respect to depth indicate that post-depositional processes did not affect the shells and therefore the <sup>87</sup>Sr/<sup>86</sup>Sr ratios are considered original.

## 3.3. Strontium isotope analyses

To determine <sup>87</sup>Sr/<sup>86</sup>Sr ratios, 0.01-0.6 mg of ostracod(s) were dissolved in 0.5 ml acetic acid and centrifuged to separate the solution from any residues. No residues were observed. The so-

lution was then pipetted into a separate beaker and evaporated to dryness and then re-dissolved in concentrated HNO<sub>3</sub> (1 drop), evaporated to dryness and again dissolved in concentrated HNO<sub>3</sub> (2 drops) and evaporated to dryness. The samples were then dissolved in 0.5 ml 3N HNO<sub>3</sub> and stored on a hotplate for 1 h. Samples were put in an ultrasonic bath for 15 min before chromatographic separation. 0.4 ml of the samples was introduced to chromatographic columns composed of "Elchrom Sr spec" ion exchange resin to isolate the strontium fraction.

Strontium isotope analyses were carried out on a Finningan MAT 262 thermal ionisation mass spectrometer (TIMS) at the Vrije University (Amsterdam) running in the 'static' mode. Strontium isotope ratios were corrected for instrumental mass fractionation by normalizing to  $^{86}$ Sr/ $^{88}$ Sr = 0.1194. Samples were analyzed over a two-year period during which degradation of Faraday cup performance led to replacement of some of the Faraday cups and changes in the cup configurations used. Data are reported for three distinct analytical periods. Repeated measurements of the international standard NBS-987 yielded averages of respectively 0.710273  $\pm$  0.000008 (2 $\sigma$ ), n = 28; 0.710259  $\pm$  0.000011 (2 $\sigma$ ), n = 18;  $0.710257 \pm 0.000006$  (2 $\sigma$ ), n = 22. Data were normalized to the long term average value obtained for NBS 987 over the previous 5 years of 0.710254  $\pm$  0.000005 (2 $\sigma$ ), n > 100. The strontium blanks were <330 pg (average 197) and negligible relative to the overall amount of strontium analyzed from the ostracods (>100 ng).

#### 3.4. Strontium isotope mass-balance models

The strontium isotope composition (<sup>87</sup>Sr/<sup>86</sup>Sr) of restricted water masses is determined by mixing of local sources such as hydrothermal vents and river water. These local inputs have their own typical isotopic composition and concentrations as a function of lithology in the catchment (Vonhof et al., 2003; Vasiliev et al., 2010). To comprehend the <sup>87</sup>Sr/<sup>86</sup>Sr of the Paratethys during the MSC, a mass-balance model was applied based on modern isotopic inputs from the most important strontium sources for these basins:

$$= \frac{(F_a \cdot ^{87/86} Sr_{Paratethys}) + (F_b \cdot ^{87/86} Sr_b) \cdots + (F_z \cdot ^{87/86} Sr_z)}{F_a + F_b \cdots + F_z}$$

wherein  $F_x$  is the strontium flux (mol/yr) and  $^{87/86}Sr_x$  is the strontium isotope composition of a certain source.

The strontium flux is given by

$$F_X = Q_X \cdot C_X$$

wherein  $F_x$  is the strontium flux (mol/yr),  $Q_x$  is the discharge (L/yr) and  $C_x$  is the strontium concentration of the water (mol/L). Note that a certain increase in discharge does not imply a proportional increase of the strontium flux, because strontium concentrations are known to drop with increasing discharge rates (Palmer and Edmond, 1989).

Since the exact isotopic compositions and flux sizes are unknown for Messinian times, we used present-day values to predict the <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the Black and Caspian Seas waters during the MSC (Table 1). We consider this approach valid because the catchments of these rivers have not changed appreciably since the late Miocene (Popov et al., 2006).

#### 3.5. *Gypsum saturation calculations*

In order to investigate the amount of evaporation necessary for gypsum formation (CaSO<sub>4</sub> \*2 H<sub>2</sub>O), calculations were performed

on modern Mediterranean and Caspian seawater using PHREEQC (Parkhurst and Appelo, 2013) at 25 °C with the Pitzer database. The chemical composition of the fluids used in the PHREEQC calculations are given in Table 2. The gypsum saturation level  $(\Omega^*_{gypsum})$  of seawater is calculated as follows:

$$\Omega_{gypsum}^* = \frac{IAP}{K_{sp}}$$

in which  $K_{sp}$  is the salinity-dependent solubility of gypsum at 25 °C (Sun et al., 2015) and IAP is the ion activity product of free dissolved calcium and sulfate,  $\{Ca^{2+}\}\{SO_4^{2-}\}$ . If  $IAP > K_{sp}$ , gypsum can precipitate from the oversaturated water. Average seawater composition (Broecker and Peng, 1982) was used for Mediterranean seawater calculation, modern Caspian seawater composition was taken from (Clauer et al., 2000) and modified Caspian seawater was obtained by removing sodium and chloride (potassium was retained and used to correct for charge imbalance). Standard evaporation (- $H_2O$ ) and mixing (MIX) options available in PHREEQC were used.

#### 4. Results

#### 4.1. Strontium isotopes

After a mechanical cleaning procedure, samples were first evaluated for potential diagenetic alteration. Samples with abundant ostracods were screened by obtaining a trace metal depth profile. Multiple trace elements were used as indicators of potential sources of contamination. Al and U are indicative of shell surface contamination by clay particles, Mn would signal secondary carbonate and/or manganese (hydr)oxides overgrowth. Although minor peaks in Al/Ca and U/Ca ratios indicated that there was still some contamination present on the outside of the ostracod shell despite the cleaning procedure (Fig. 3), the quantity of these 'contamination peaks' is low in comparison to the total quantity of the inner shell. For example, <sup>88</sup>Sr counts in the <sup>24</sup>Mg/<sup>44</sup>Ca peak of AJ79 account only for <0.1% of the Sr content of the shell. The constant elemental ratios in the ablation profiles furthermore indicate that post-depositional processes such as recrystallization and/or solidstate diffusion did not affect <sup>87</sup>Sr/<sup>86</sup>Sr and therefore none of the samples were considered appreciably affected by diagenetic alteration.

Our strontium isotope records from the Paratethys indicate three main hydrological phases in the late Miocene (Fig. 2; Table 3). Predating the MSC, <sup>87</sup>Sr/<sup>86</sup>Sr ratios in the Paratethys are relatively low (0.70825) and close to the present-day Caspian Sea values (0.7082) (Clauer et al., 2000). At present, the Caspian Basin is isolated and its <sup>87</sup>Sr/<sup>86</sup>Sr ratio is mainly determined by the Volga river and hydrothermal vents that supply relatively unradiogenic Sr (Clauer et al., 2000). The similarity in <sup>87</sup>Sr/<sup>86</sup>Sr ratios with the present-day Caspian Sea indicates that the pre-MSC Paratethys Sea was disconnected from the ocean as well. At  $\sim$ 6.1 Ma, a rise in 87Sr/86Sr ratios is recorded, with the Caspian Sea data approaching open marine values (Fig. 2). The increase in <sup>87</sup>Sr/<sup>86</sup>Sr ratios coincides with the regional Pontian Flood (Krijgsman et al., 2010; Grothe et al., 2014; Van Baak et al., 2017). This marine transgressive event connected the Black and Caspian Seas with the Mediterranean and some central European (Dacian and Pannonian) basins (Popov et al., 2006). Remarkably, marine ratios are not recorded in our Black Sea section, possibly because bottom water anoxia prevented the presence of ostracods during the 'peak-flooding'. Following this transgressive event, 87Sr/86Sr-values of the Black and Caspian Seas decrease again and cluster around 0.7084-0.7085 throughout the MSC interval (Fig. 2). Although the Caspian Sea  $^{87}$ Sr/ $^{86}$ Sr ratios are generally slightly lower ( $\sim$ 0.7084), the similarity with the Black Sea values strongly supports a sustained

**Table 2**Chemical composition of the fluids used in the PHREEQC calculations: Mediterranean Seawater (Med. SW; after Nordstrom et al., 1979), Caspian/Black Seawater (C/B SW, after Clauer et al., 2000 except for Dissolved Inorganic Carbon (DIC), which is from Meybeck, 1993), younger C/B SW (yC/B SW; without sodium chloride) and C/B SW and yC/B SW evaporated until equilibrium with gypsum was reached (with subscript evap).

|                        | Med. SW        | C/B SW   | C/B SW <sub>evap</sub> | yC/B SW  | yC/B SW <sub>evap</sub> |
|------------------------|----------------|----------|------------------------|----------|-------------------------|
| pН                     | 8.2            | 7.0      | 6.9                    | 7.0      | 7.0                     |
| Salinity               | 37             | 15       | 43                     | 15       | 14                      |
| $\Omega_{gypsum}$      | 0.20           | 0.33     | 1.00                   | 0.33     | 1.00                    |
| Total concentration    | ons (mmol/kgw) |          |                        |          |                         |
| DIC                    | 2.20           | 0.44     | 1.24                   | 0.44     | 0.80                    |
| Total Ca               | 10.9           | 8.03     | 22.7                   | 8.03     | 14.6                    |
| Total Cl               | 575            | 143      | 405                    | 0        | 0                       |
| Total K                | 11.0           | 2.12     | 6.00                   | 2.12     | 3.87                    |
| Total Mg               | 55.1           | 28.8     | 81.3                   | 28.8     | 52.5                    |
| Total Na               | 494            | 131      | 369                    | 0        | 0                       |
| Total SO <sub>4</sub>  | 29.8           | 31.2     | 88.3                   | 31.2     | 57.0                    |
| Ion (pair) activitie   | es (unitless)  |          |                        |          |                         |
| $\{HCO_3^-\}$          | 1.19E-03       | 2.72E-04 | 6.30E-04               | 2.73E-04 | 4.63E-04                |
| $\{CO_3^{2-1}\}$       | 9.01E-06       | 1.24E-07 | 2.20E-07               | 1.25E-07 | 1.92E-07                |
| $\{Ca^{2+}\}$          | 2.47E-03       | 1.87E-03 | 4.16E-03               | 2.07E-03 | 3.06E-03                |
| {Cl <sup>-</sup> }     | 3.58E-01       | 9.95E-02 | 2.51E-01               | 0        | 0                       |
| $\{K^+\}$              | 7.16E-03       | 1.50E-03 | 3.84E-03               | 1.63E-03 | 2.84E-03                |
| ${Mg^{2+}}$            | 1.37E-02       | 7.52E-03 | 1.73E-02               | 8.74E-03 | 1.31E-02                |
| $\{MgCO_3^0\}$         | 1.05E-04       | 7.93E-07 | 3.23E-06               | 9.24E-07 | 2.13E-06                |
| {MgOH <sup>+</sup> }   | 3.47E-06       | 1.16E-07 | 2.02E-07               | 1.36E-07 | 1.84E-07                |
| {Na <sup>+</sup> }     | 3.49E-01       | 9.55E-02 | 2.55E-01               | 0        | 0                       |
| ${SO_4^{2-}}$          | 2.49E-03       | 5.25E-03 | 7.38E-03               | 7.26E-03 | 9.83E-03                |
| Ion activity ratios    | i              |          |                        |          |                         |
| ${Ca^{2+}}/{Cl^{-}}$   | 6.88E-03       | 1.88E-02 | 1.66E-02               |          |                         |
| ${SO_4^{2-}}/{Cl^{-}}$ | 6.94E-03       | 5.28E-02 | 2.94E-02               |          |                         |

connection between these two basins. We have no data from the Black Sea between 5.6-5.3 Ma because of a stratigraphic hiatus in the section, but the stability of the Caspian Sea values, as well as biogeographic data, suggests that these basins remained connected (Vasiliev et al., 2013).

#### 4.2. Mass balance model for the Paratethys Sea

The  $^{87}$ Sr/ $^{86}$ Sr of the Paratethys seawater was calculated for four different input scenarios (Table 1):

- (i) main rivers only (Danube, Dnieper Don, Volga);
- (ii) main rivers and hydrothermal vents;
- (iii) main rivers, hydrothermal vents and Mediterranean inflow;
- (iv) main rivers and Mediterranean inflow.

We consider the clustering of <sup>87</sup>Sr/<sup>86</sup>Sr ratios around 0.7084-0.7085 to represent a system in which the Black and Caspian Seas are 'fully connected'. Mass balance calculations for scenarios (i) and (ii), based on modern strontium input from the most important sources, however, yields a lower  $^{87}$ Sr/ $^{86}$ Sr ratio:  $\sim$ 0.7083 and 0.7082, respectively. Therefore, a source of more radiogenic Sr would be required to explain our record. Since the late Miocene drainage system was very similar to present-day (Popov et al., 2006), the most likely additional source for high <sup>87</sup>Sr/<sup>86</sup>Sr ratios is open marine water. To quantify how much inflow of marine waters is required to explain our measured strontium records, scenario (iii) and (iv) were evaluated. Adding an open marine source via a "Bosphorus"-like strait requires only ~4-10% of present-day marine input to explain the observed Sr isotope data for the Black and Caspian Seas (scenario iii). A further reduction of the presentday inflow down to 1-5% is enough to approach the measured <sup>87</sup>Sr/<sup>86</sup>Sr ratios with scenario (iv) in the mass-balance. Such an inferred marine inflow into the Black Sea agrees with late Miocene proxy records showing presence of alkenones typical for marine conditions in the Black Sea (Vasiliev et al., 2013).

#### 5. Discussion

# 5.1. Paratethys pacing of the MSC

Combining our new Black and Caspian Sea Sr isotope records with that of the Mediterranean, four main circulation modes can be identified between 6.7 and 5.0 Ma (Fig. 2).

be identified between 6.7 and 5.0 Ma (Fig. 2).

<u>Mode 1 (6.7-6.1 Ma)</u>: Before 6.1 Ma, <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the Black and Caspian Seas are the lowest. The similarity in <sup>87</sup>Sr/<sup>86</sup>Sr ratios with the present-day isolated Caspian Sea (Fig. 2), coupled with our mass balance calculations, indicates that the two basins were disconnected from the open ocean forming a unified Paratethys Lake, which was dominantly filled by local rivers. Coeval <sup>87</sup>Sr/<sup>86</sup>Sr values from the Mediterranean mostly plot on the ocean curve, attesting that the Mediterranean was a marine basin connected to the Atlantic.

Mode 2 (6.1-5.55 Ma): At  $\sim$ 6.1 Ma, a major transgressive event (i.e. the Pontian Flood after Krijgsman et al., 2010) connected the Paratethys to the Mediterranean. Despite the low resolution record, this short-lived event is reflected in the Paratethys Sr signal which shows a peak at 6.1 Ma, reaching values close to the marine curve (Fig. 2). The inflow of marine water into the fresh-brackish Paratethys lake may have created a two-way exchange at the gateway (cf. Marzocchi et al., 2016). The establishment of a Paratethyan flow towards the Mediterranean has recently been detected in the biomarker records of the Eastern Mediterranean that show a sudden freshening of the surface waters at  $\sim$ 6.1 Ma (Vasiliev et al., 2019).

The <sup>87</sup>Sr/<sup>86</sup>Sr values of MSC Stage 1 and 2 evaporites are significantly offset from the oceanic curve (Fig. 2), indicating that large amounts of non-marine water spilled into the Mediterranean. It is not surprising that Mediterranean <sup>87</sup>Sr/<sup>86</sup>Sr measured on the MSC Primary Lower Gypsum (Stage 1) remain close to oceanic values because the connection with the Atlantic Ocean, although restricted, was still rather efficient. The onset of the MSC at 5.97 Ma is often linked to the time that <sup>87</sup>Sr/<sup>86</sup>Sr of Mediterranean

**Table 3** Sample codes, stratigraphic position, regional stages, ages (in Ma) levels, ostracod species and  $^{87}$ Sr $^{186}$ Sr ratios with  $2\sigma$  error of all analysed samples.

| Sample | le Stratigraphic Paratethys (sub)stages<br>level<br>(m) |                        | Age<br>(Ma) | Rough age<br>estimate<br>(Ma) | Ostracods and foraminifera species          | <sup>88</sup> Sr/ <sup>87</sup> Sr |
|--------|---|------------------------|-------------|-------------------------------|---|------------------------------------|
| TR23   | -329.00   | Upper Maeotian         | 6.2 - 8.2   | 7.75                          | Cyprideis (juveniles)                       | $0.708084 \pm 10$                  |
| TH36   | -298.00   | Upper Maeotian         | 6.2 - 8.2   | 7.2                           | Condon? neglecta                            | $0.708234 \pm 9$                   |
| TR27   | -269.00   | Upper Maeotian         | 6.2 - 8.2   | 6.7                           | Ammonia beccom                              | $0.708228\pm10$                    |
| TR30   | -266.00   | Upper Maeotian         | 6.2 - 8.2   | 6.6                           | Cyprideis sp.                               | $0.708229 \pm 8$                   |
| TR41   | -255.00   | Upper Maeotian         | 6.2 - 8.2   | 6.55                          | Ammonia beccom $+$ Cyprideus sp. (juvenile) | $0.708286 \pm 8$                   |
| TR71   | -206.00   | Upper Maeotian         | 6.160       | -                             | Cyprideis sp.                               | $0.708284 \pm 16$                  |
| TR80   | -197.00   | Pontian Foram interval | 6.100       | -                             | Cyprideis, Leptocyther sulakensis           | $0.708234 \pm 7$                   |
| TR80   | -197.00   | Pontian Foram interval | 6.100       | -                             | Cyprideis sp.                               | $0.708231\pm10$                    |
| TR82   | -195.00   | Pontian Foram interval | 6.090       | -                             | Cyprideis                                   | $0.708265 \pm 8$                   |
| TR83-c | -194.00   | Pontian Foram interval | 6.088       | -                             | Cyprideis sp.                               | $0.708220 \pm 9$                   |
| TR89   | -185.00   | Pontian                | 6.050       | -                             | Pontoniella lotzyr                          | $0.708397 \pm 9$                   |
| TR89a  | -185.00   | Pontian                | 6.050       | -                             | Amnicythere sp.                             | $0.708433\pm10$                    |
| TR92   | -182.00   | Pontian                | 6.040       | -                             | Cyprideis sp.                               | $0.708400 \pm 9$                   |
| TR101  | -173.00   | Pontian                | 6.000       | -                             | Caspiocypris atta                           | $0.708404 \pm 10$                  |
| TR109  | -165.00   | Pontian                | 5.960       | -                             | Caspiocypris atta                           | $0.708415 \pm 9$                   |
| TR115  | -148.00   | Pontian                | 5.900       | -                             | Caspiocypris atta                           | $0.708503 \pm 16$                  |
| TR117  | -146.00   | Pontian                | 5.870       | -                             | Caspiocypris atta                           | $0.708474 \pm 9$                   |
| TR126  | -133.00   | Pontian                | 5.840       | -                             | Caspiocypris atta                           | $0.708531 \pm 9$                   |
| TK38   | -117.00   | Pontian                | 5.750       | _                             | Caspiocypris atta                           | $0.708505 \pm 9$                   |
| TK24   | -107.00   | Pontian                | 5.670       | _                             | Caspiocypris atta                           | $0.708567 \pm 9$                   |
| TK18   | -101.00   | Pontian                | 5.630       | -                             | Caspiocypris atta                           | $0.708538 \pm 9$                   |
| TK12   | -96.00  | Pontian                | 5.620       | -                             | Caspiocypris atta                           | $0.708520 \pm 9$                   |
| TK09   | -94.00  | Pontian                | 5.610       | -                             | Caspiocypris? atta                          | $0.708608 \pm 9$                   |
| TK52   | -77.00  | Kimmerian              | 5.100       | -                             | Casprocypiris (fragments)                   | $0.708434 \pm 9$                   |
| AA18   | 5.60  | Upper Maeotian         | 6.125       | -                             | Leptocythere sulakensis                     | $0.708297\pm8$                     |
| AA22   | 7.00  | Pont: Foram interval   | 6.110       | -                             | Leptocythere sulakensis                     | $0.708729 \pm 10$                  |
| AA24   | 7.58  | Pont: Foram interval   | 6.080       | -                             | Leptocythere sulakensis                     | $0.708331 \pm 9$                   |
| AA114  | 33.60   | Pontian                | 5.930       | -                             | Amnicythere palimpsesta                     | $0.708360 \pm 9$                   |
| AA134  | 39.80   | Pontian                | 5.870       | -                             | Amnicythere palimpsesta                     | $0.708406 \pm 12$                  |
| AA146  | 42.90   | Pontian                | 5.850       | -                             | Amnicythere palimpsesta, Caspiocypris       | $0.708277 \pm 10$                  |
| AA150  | 43.95   | Pontian                | 5.840       | -                             | Amnicythere palimpsesta                     | $0.708277 \pm 9$                   |
| AA180  | 51.75   | Pontian                | 5.790       | -                             | Amnicythere palimpsesta                     | $0.708422\pm8$                     |
| AA200  | 57.00   | Pontian                | 5.760       | -                             | Amnicythere palimpsesta                     | $0.708423 \pm 9$                   |
| AJ09   | 59.00   | Pontian                | 5.750       | -                             | Amnicythere palimpsesta                     | $0.708488 \pm 9$                   |
| AJ19   | 61.50   | Pontian                | 5.740       | -                             | Amnicythere palimpsesta                     | $0.708524 \pm 9$                   |
| AA215  | 61.55   | Pontian                | 5.740       | -                             | Amnicythere palimpsesta                     | $0.708494 \pm 9$                   |
| AJ26   | 64.00   | Pontian                | 5.730       | -                             | Amnicythere palimpsesta                     | $0.708457 \pm 9$                   |
| AJ79   | 76.75   | Pontian                | 5.630       | -                             | Amnicythere palimpsesta                     | $0.708485 \pm 9$                   |
| AJ111  | 84.75   | Pontian                | 5.590       | -                             | Amnicythere palimpsesta                     | $0.708442 \pm 9$                   |
| AJ123  | 87.75   | Pontian                | 5.580       | _                             | Amnicythere palimpsesta                     | $0.708435 \pm 13$                  |
| AJ142  | 92.50   | Pontian                | 5.550       | -                             | Amnicythere palimpsesta                     | $0.708402 \pm 9$                   |
| AJ204  | 113.50  | Pontian                | 5.450       | _                             | Amnicythere palimpsesta                     | $0.708433 \pm 7$                   |
| AJ217  | 119.75  | Pontian                | 5.420       | _                             | Amnicythere palimpsesta                     | $0.708456 \pm 8$                   |
| AJ223  | 122.00  | Pontian                | 5.410       | _                             | Amnicythere palimpsesta                     | $0.708452\pm10$                    |
| AJ234  | 127.75  | Pontian                | 5.380       | -                             | Amnicythere palimpsesta                     | $0.708473 \pm 9$                   |

seawater starts to deviate from open ocean Sr-isotope values, but <sup>87</sup>Sr/<sup>86</sup>Sr of foraminifers just preceding the onset of the MSC already show an offset (Fig. 2; Roveri et al., 2014b). Strontium isotope data from individual gypsum beds of the Northern Apennines show that gypsum precipitation started from a seawater body marked by non-marine contributions and that the influence of marine input increased through time within a precessional cycle (Reghizzi et al., 2018). These Sr-isotope values are so far ascribed to variations in freshwater discharge from the major peri-Mediterranean rivers only (e.g. Rhône and Nile) (García-Veigas et al., 2018; Reghizzi et al., 2018). We postulate here that Paratethys may have contributed to the freshening of the Mediterranean surface waters during PLG formation. The westward spread of brackish Paratethyan waters could have been facilitated by the density contrast with the saline marine Mediterranean, which prevents mixing and favors a strongly stratified water column (cf. the MSC scenario of García-Veigas et al., 2018). The absence of both Paratethyan and Atlantic faunal assemblages in MSC Stage 1 and 2 deposits suggests that an ecological barrier (probably salinity) was present that prevented faunal exchange with the Mediterranean during this time.

Mode 3 (5.55-5.33): The low  ${}^{87}$ Sr/ ${}^{86}$ Sr values of MSC Stage 3 (0.7085-0.7087) imply a major contribution from Paratethyan wa-

ters into the Mediterranean, at least within the surface layer, and a further reduction of Atlantic inflow (Fig. 2). To approach the <sup>87</sup>Sr/<sup>86</sup>Sr values that characterize Stage 3, the present-day inflow of the Atlantic needs to be reduced by at least a factor of 1000. With a strongly reduced Atlantic inflow, a volume of Paratethys Sea inflow similar to present-day Black Sea inflow is sufficient to approach the observed Mediterranean <sup>87</sup>Sr/<sup>86</sup>Sr. A strongly reduced Atlantic inflow, however, would lead to a drop in Mediterranean water levels, which is in line with the classic desiccation model (Hsü et al., 1973), but not with the idea that the Mediterranean remained water-filled (Roveri et al., 2014a; Stoica et al., 2016). Reduced mixing between a low salinity surface water layer and deeper highly saline Atlantic waters provides an alternative mechanism (Marzocchi et al., 2016; García-Veigas et al., 2018). The high density of the brine responsible for halite precipitation (MSC Stage 2) may have prevented waters from the Paratethys to mix due to the large density contrast. Consequently, reduced mixing of these low salinity Paratethyan waters would facilitate the spread of these waters as a thin but extensive low salinity lid across the entire Mediterranean. Environmental conditions in the Mediterranean during Upper Gypsum deposition (MSC substage 3.1) were still unfavorable for most Paratethyan fauna to migrate westward.

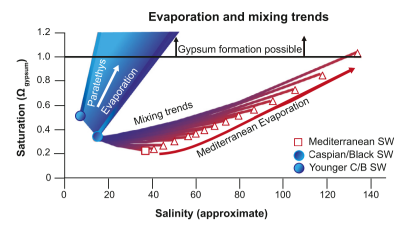


Fig. 4. Model results for evaporation of Mediterranean and Caspian seawater during the MSC. Square is for average seawater (SW) and triangles for progressive evaporation of this SW; light circle is for modern Caspian SW (Clauer et al., 2000) and light diamonds for progressive evaporation of this water; dark circle is for modified Caspian SW (sodium chloride free, i.e., NaCl-free) and light diamonds for progressive evaporation of modified Caspian SW; the horizontal solid line indicates equilibrium with gypsum and above this line, gypsum precipitation is thermodynamically possible; other solid lines relate to mixing of (evaporated) Mediterranean SW and modern Caspian SW.

A marked hydrological change took place in the Mediterranean at 5.42 Ma and the onset of the Lago-mare (MSC substage 3.2; Roveri et al., 2014a), when numerous Paratethyan ostracod and mollusk species invaded the Mediterranean (Gliozzi et al., 2007; Stoica et al., 2016). It must be noted here that also during substage 3.2 some gypsum layers formed in the Mediterranean (e.g., Eraclea Minoa; Roveri et al., 2014a). The density contrast between Mediterranean and Paratethyan waters may have triggered a huge outflow pump (Marzocchi et al., 2016) that can explain the existence of Black Sea endemic fauna in even the westernmost Mediterranean basins during MSC Stage 3.2 (Guerra-Merchán et al., 2010; Stoica et al., 2016).

Increased input from the Nile River or other major peri-Mediterranean rivers might have added to the observed isotopic ratios (Gladstone et al., 2007), but evidence exists for inflow from the Black Sea in the form of typical Paratethys fauna (Gliozzi et al., 2007; Stoica et al., 2016). We postulate here that inflow from the Paratethys forms an alternative and simpler scenario to explain the low <sup>87</sup>Sr/<sup>86</sup>Sr values of Stage 3. The extensive hinterland draining into the Black and Caspian Sea (Fig. 1) delivers water roughly one order of magnitude higher in Sr concentrations compared to Nile river water. Hence, although the Nile might also have added low radiogenic strontium, the observed temporal changes in <sup>87</sup>Sr/<sup>86</sup>Sr are alternatively explained by a Paratethys source. Drainage from other North African basins (including Lake Chad) would have, in contrast to the Nile, increased the 87Sr/86Sr ratios of the Mediterranean because their catchments lack extensive Rbdepleted basaltic rocks with a characteristic low 87 Sr/86 Sr (Topper et al., 2014).

Mode 4 (5.33-5.0 Ma): Finally, during the onset of the Pliocene, the Atlantic-Mediterranean connection is restored and marine conditions return in the Mediterranean. Paratethys most likely continued to drain low salinity water into the Mediterranean, but this is no longer detected in the strontium isotope values. The restored connection to the Atlantic Ocean converted Mediterranean <sup>87</sup>Sr/<sup>86</sup>Sr ratios towards oceanic values, because ocean water has a 7-8 times higher Sr concentration than non-marine sources.

#### 5.2. A brackish water trigger for gypsum formation during the MSC?

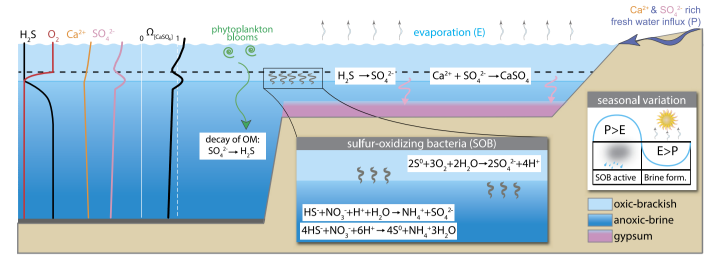
The new strontium isotope data strongly suggest the existence of a sustained connection between the Mediterranean and Paratethys during the entire MSC. This connection would have affected the Mediterranean water budget by adding low salinity water. While restriction of the Mediterranean-Atlantic gateway(s) is considered the principal criterion in explaining the formation of evaporites (Garcia-Castellanos and Villaseñor, 2011; Flecker et al.,

2015; Krijgsman et al., 2018), our data suggest that the onset of a Mediterranean-Paratethys connection, which preceded the first gypsum beds, somehow played a major role in shaping the MSC-events.

The gypsum deposits of MSC Stage 1 are all from shallowwater settings (<200 m) and contain low-salinity fluid inclusions with stable oxygen isotope signals indicative of large inputs of Na<sup>+</sup> and Cl<sup>-</sup>-depleted waters during their formation (Natalicchio et al., 2014; Evans et al., 2015). Natalicchio et al. (2014) proposed that local rivers enriched in  $Ca^{2+}$  and  $SO_4^{2-}$ , derived from partial dissolution and recycling of coeval marginal marine deposits, are the main source of non-marine  $Ca^{2+}$  and  $SO_4^{2-}$ . We here propose Paratethys may have contributed in three ways: 1) Freshwater from the Paratethys entering the strongly restricted Mediterranean Basin would certainly enhance stratification, resulting in a low-salinity surface-water layer that could influence the water composition in the individual shallow basins where MSC Stage 1 gypsum accumulated; 2) the freshwater surface-water from the Paratethys had relatively high concentrations of  $Ca^{2+}$  and  $SO_4^{2-}$  (cf. Clauer et al., 2000), and may have added  $Ca^{2+}$  and  $SO_4^{2-}$  to the underlying seawater layer resulting in gypsum precipitating from a marine layer; 3) the low-salinity water with relatively high concentrations of  $Ca^{2+}$  and  $SO_4^{2-}$  itself provided the non-marine setting in which gypsum could precipitate by evaporation.

Interestingly, the Paratethyan waters flowing in to the Mediterranean are as high in calcium and sulfate as average seawater (cf. Clauer et al., 2000). At the same time, Paratethyan water has much lower sodium and chloride concentrations than average seawater (Table 2). These ions play a crucial role in gypsum formation, by affecting the ion activity product (IAP,  $\{Ca^{2+}\}\{SO_4^{2-}\}$ ; see section 3.4). Dissolved sodium and chloride pair with sulfate and calcium in solution, respectively, thereby lowering the IAP and the saturation state ( $\Omega_{gypsum}^*$ ). In other words, high chloride and sodium concentrations, as found in seawater and evaporated seawater, increase gypsum solubility, and make it harder to reach super-saturated conditions in which gypsum can precipitate.

The impact of sodium and chloride concentrations on saturation state is clear: while average seawater and Caspian water have similar total calcium and sulfate concentrations, seawater is more strongly undersaturated (Fig. 4; Table 2). Moreover, to reach gypsum saturation normal seawater has to be concentrated by evaporation to a salinity of  $\sim$ 120. In contrast, evaporation of modern Caspian water becomes gypsum saturated at a salinity of  $\sim$ 40 (Fig. 4). Mixing Mediterranean Sea-water, or any of the solutions along its evaporation trend, with modern Caspian water enhances saturation but not enough for gypsum precipitation.



**Fig. 5.** Conceptual model illustrating the suggested role of sulfur-oxidizing bacteria (SOB) in biogeochemically meditated gypsum formation. SOB inhabiting a niche on the pycnocline profit from the availability of sulfide and oxygen. SOB transform reduced sulfur species  $(H_2S, S_0)$  into sulfate  $(SO_4^{2-})$  and thereby raise the production of  $[Ca^{2+}]$  and  $[SO_4^{2-}]$ , which may trigger gypsum precipitation. The nutrients in the upper part of the water-column promote productivity. Subsequent decay of this organic material in anoxic waters produces sulfides  $(H_2S)$ , which are required to sustain the SOB-activity.

Miocene Paratethyan waters may even have had lower sodium and chloride contents than present-day Caspian water, because Paratethys was not the closed evaporative basin as the Caspian has been over the last  $\pm 3$  Ma (Palcu et al., 2019). High concentrations of sodium and chloride in sea water are the result of billions of years of weathering and the long residence times of these elements and consequently young waters are relatively impoverished in both elements. Assuming calcium and sulfate concentrations similar to modern Caspian water and negligible sodium and chloride, then evaporation of Paratethyan water leads to gypsum saturation at a salinity of  $\pm 14$  (Fig. 4). If the Paratethys water was lower in calcium and sulfate then assumed here, the saturation state would be lower (  $\Omega^*_{\it gypsum}$  would be smaller), but the evaporation trend would be as steep as plotted in Fig. 4, leading to a similarly low salinity for gypsum saturation. The product of  $\{Ca^{2+}\}\{SO_{\Delta}^{2-}\}\$  can also rise due to an increased influx of calcium and/or sulfate. Two superimposed water layers, both enriched in sulfate and calcium, can exchange ions by diffusion and influence the gypsum solubility product without increasing the overall salinity (Fig. 5). Additional sulfate can be provided by abiotic (e.g. Lin et al., 2016) or biologic (e.g. Bailey et al., 2009) oxidation of reduced sulfur compounds. The MSC gypsum and shale deposits contain ubiquitous large filamentous microfossils, interpreted initially as remains of algae (Vai and Ricci Lucchi, 1977), faecal pellets (Guido et al., 2007) or cyanobacteria (Panieri et al., 2010). More recently they have been interpreted as remains of sulfide-oxidizing bacteria (SOB), based on the identification of sulfur-containing minerals possibly derived from the diagenetic transformation of elemental sulfur stored within cytoplasm (Dela Pierre et al., 2014, 2015; Andreetto et al., 2019). SOBs do not provide useful paleobathymetrical and salinity information, but they play a key role in the biogeochemical sulfur cycle by producing  ${\rm SO_4^{2-}}$  from the oxidation of reduced sulfur species [e.g. H<sub>2</sub>S, S<sup>2-</sup>] (e.g. Bailey et al., 2009). The exact role of microbes in gypsum precipitation is still poorly understood, but potential additional sulfate input from SOB would result in super saturation being reached at lower levels of evaporation than shown in Fig. 4 and hence potential gypsum formation at even lower salinities (Fig. 5).

#### 5.3. Paratethys contribution to other gypsum deposits

Our observations suggest that shallow-basin MSC gypsum layers can be formed by evaporation of a low salinity water layer,

partly sourced from the Paratethys Sea. Gypsum formation being influenced by Paratethyan water was probably not restricted to the Mediterranean MSC as numerous gypsum deposits are known from several former Paratethyan basins. Archetypes are the evaporites of the Badenian Salinity Crisis (13.8-13.4 Ma; Peryt, 2006; De Leeuw et al., 2010) of the Central Paratethys (Carpathian Foredeep, Transylvanian Basin, Pannonian Basin), and several other gypsum units have been described from the early and middle Miocene Paratethyan basins as well (see De Leeuw et al., 2018 and references therein). The paleogeographic configuration of the Badenian and Messinian Salinity Crises are very similar, with the evaporite basins in intermediate position between the open ocean and the restricted brackish water domain of the Eastern Paratethys (Palcu et al., 2017). More detailed geochemical studies of the Badenian and Messinian evaporites are needed to better understand the role of Eastern Paratethys waters, and the potential involvement of SOB on gypsum formation.

#### 5.4. Pitfalls and future perspectives

All data so far indicate that Paratethys-Mediterranean connectivity existed during the latest Messinian and that Paratethyan water, with much higher concentrations of strontium, calcium and sulfate than river water, contributed as an additional low salinity water source to the Mediterranean hydrological balance during the different MSC stages. The remaining key question is now: How significant was the late Messinian Paratethys flux of low salinity water for the Mediterranean? And should the various hydrological. strontium, gypsum and climate models of the MSC be updated and revised? At this moment, no quantitative amounts of Paratethyan water fluxes are available for MSC times. In addition, the chemical composition of the Messinian Paratethys is still unknown; in this paper we have used the present-day Caspian values as the most likely modern analogue. More detailed studies on the paleoenvironmental and paleohydrological changes that occurred in Paratethys during the Messinian (Pontian stage in regional terminology) are therefore very welcome.

Despite the uncertainties in the exact location of the gateway(s) (see Van Baak et al., 2016), Mediterranean-Paratethys connectivity most likely has taken place via the Aegean domain (Fig. 1), an area with poorly dated Miocene successions that is generally considered as an intermediate domain, showing aspects of both Mediterranean and Paratethyan fauna (Popov et al., 2006).

Paratethyan fauna is absent in the Mediterranean during MSC Stage 1 and 2, sporadically occurring in MSC substage 3.1 (Eraclea Minoa, Sicily; Grossi et al., 2015), but omnipresent in substage 3.2. This trend may indicate an increase through time of the relative contribution of Paratethyan waters to the Mediterranean and/or a progressive change in Mediterranean salinity, which is the one of the main environmental factor that controls the proliferation of brackish Paratethyan organisms during the MSC. This salinity change is most likely amplified by the significant decrease, or even complete termination, of the Atlantic influx through the Gibraltar Corridor during Stage 3 (see Krijgsman et al., 2018 for details). Revised hydrological and climate models are thus necessary to test for different Atlantic-river-Paratethys influx values on Mediterranean strontium ratios and gypsum precipitation models. This will eventually also solve the question to what stage(s) the Paratethys has influenced the Mediterranean MSC.

#### 6. Conclusions

Strontium isotope records from the Paratethys reveal several hydrological changes in the latest Miocene. Predating the MSC (>6.1 Ma),  $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$  ratios are relatively low (0.70825), indicating that the Paratethys was mainly disconnected from the Mediterranean. At 6.1 Ma, a sudden rise in  $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$  (max. 0.7087) corresponds to the Pontian Flood and reflects the initiation of Paratethys-Mediterranean connectivity. Throughout the MSC interval (5.97-5.33 Ma),  $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$  ratios in Paratethys cluster around 0.7084-0.7085, values that are best explained by a severely reduced connection with <10% of the present-day inflow through the Bosphorus Strait.

The strontium isotope variations of Paratethys suggest a direct pacing of Mediterranean hydrology and MSC gypsum formation. The Mediterranean  $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$  record shows a deviation from the open ocean curve at 6.1 Ma, when a connection with the Paratethys is established (Fig. 2). During MSC Stage 3 the Mediterranean  $^{87}\mathrm{Sr}/^{86}\mathrm{Sr}$  values (down to  $\sim\!0.7085$ ) appear dominated by Paratethys waters, combined with a strongly reduced contribution of the Atlantic.

We propose that the high-density brine formation during the Mediterranean MSC may have prevented mixing of Paratethys waters forming an extensive low-salinity surface layer rich in calcium and sulfate. We show that evaporation of such a surface layer results in saturation with respect to gypsum at a salinity of  $\sim$ 40. This is in good agreement with recent salinity reconstructions of MSC gypsum based on fluid inclusions (Natalicchio et al., 2014; Evans et al., 2015).

This proposed mechanistic link between gypsum formation and low-saline water inputs from the Black Sea implies that climatic phenomena and evaporation rates in the Paratethys region, and the Atlantic Ocean, should be considered as additional potential drivers for the cyclic gypsum beds and low <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the Mediterranean MSC rather than African monsoon driven input from the Nile only. Our scenario provides new constraints for future quantitative models, that should include a supplementary Paratethys component in the Mediterranean hydrological balance and assess the proportion of riverine and Paratethys <sup>87</sup>Sr/<sup>86</sup>Sr to lower the <sup>87</sup>Sr/<sup>86</sup>Sr of the MSC Mediterranean water mass during gypsum precipitation.

#### **Author contributions**

AG, GJR, CvB, IV, FS, WK all contributed to the conceptualization of the research. AG, CvB, IV, MS, WK sampled the sections, MS, FS were in charge of the faunal analyses, AG, GJR, GD performed and analyzed the strontium measurements, GJR, MW, JJM performed and analyzed the geochemical models, AG, GJR, FS, WK prepared

the original draft, FA, MW, GJR, IV, GD and WK took care of the revisions and editing of the final manuscript.

#### **Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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