# **Barium in seawater**

# Dissolved distribution, relationship to silicon, and barite saturation state determined using machine learning

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# Abstract

Barium is widely used as a proxy for dissolved nutrients silicon and particulate organic carbon fluxes in seawater. However, these proxy applications are limited by insufficient knowledge of the dissolved distribution of Ba ([Ba]). For example, there is significant spatial variability in the Ba-Sibarium-silicon relationship, and ocean chemistry may influence sedimentary Ba preservation. To help address these issues, we developed 4,095 models for predicting [Ba] using Gaussian Progress Regression Machine Learning. These models were trained to predict [Ba] from standard oceanographic observations using GEOTRACES data from the Arctic, Atlantic, Pacific, and Southern Oceans. Trained models were then validated by comparing predictions against withheld [Ba] data from the Indian Ocean. We find that a model trained using depth,  $T_{...,S_{i}}$  [O<sub>2</sub>]. [PO<sub>4</sub>], temperature, salinity, as well as dissolved dioxygen, phosphate, nitrate, and [NO<sub>3</sub>] as predictors silicate can accurately predict [Ba] in the Indian Ocean with a mean absolute percentage deviation of 6.30 %. We use this model to simulate [Ba] on a global basis using these same sixseven predictors in the World Ocean Atlas. The resulting [Ba] distribution constrains the total Ba budget of the ocean to  $122\pm8\times(\pm7)\times10^{12}$  mol and clarifies reveals systematic variability in the globalbarium-silicon relationship between dissolved Ba and Si. We also calculate the saturation state of seawater with respect to barite. In addition to revealing systematic spatial and vertical variations, our results show that the ocean below 1,000 m is, on average, at or near saturation equilibrium with respect to barite. We describe a number of possible applications for our model output, ranging from use in biogeochemical models to paleoproxy calibration. Our approach could be extended to other trace elements with relatively minor adjustments and demonstrates the utility of machine learning to accurately simulate the distributions of tracers in the sea and provides a framework that could be extended to other trace elements.

# 1. Introduction

Barium (Ba) is a Group II trace metal that is widely applied in studies of modern and ancient marine biogeochemistry, despite lacking a recognized biochemical function (e.g., Horner & Crockford, 2021). These applications of Ba are based on two empirical correlations relating to its dissolved and particulate cycles. The first correlation relates to the dissolved concentration of Ba, hereafter [Ba], which is strongly correlated with that of the algal nutrient silicon (Si; as dissolved silicic acid; Fig. 1; Chan et al., 1977). Unlike [Si], ambient [Ba] concentrations are faithfully recorded by a number of marine carbonates, such as planktonic (e.g., Hönisch et al., 2011) and benthic foraminifera (e.g., Lea & Boyle, 1990), surface- (e.g., Gonneea et al., 2017) and deep-sea corals (e.g., Anagnostou et al., 2011; LaVigne et al., 2011), and mollusks (e.g., Komagoe et al., 2018). Preservation of these signals means that the Ba content of carbonates can be related to the Ba content of seawater and, by extension, that of Si. Accordingly, the Ba–Si proxy has been applied to understand ocean nutrient dynamics on decadal (e.g., Lea et al., 1989) to millennial timescales (e.g., Stewart et al., 2021).

The nutrient-like distribution of dissolved Ba in seawater is thought to be sustained by the second empirical correlation, relating to cycling of particulate Ba. Particulate Ba in seawater occurs mostly in the form of discrete, micron-sized crystals of the mineral barite (BaSO<sub>4</sub>(s), barium sulfate; e.g., Dehairs et al., 1980; Stroobants et al., 1991). Pelagic BaSO<sub>4</sub> is an ubiquitous component of marine particulate matter (e.g., Light & Norris, 2021) and constitutes the principal removal flux of dissolved Ba from seawater (Paytan & Kastner, 1996). Pelagic BaSO<sub>4</sub> is thought to precipitate within ephemeral particle-associated microenvironments that develop during the microbial oxidation of sinking organic matter (e.g., Chow & Goldberg, 1960; Bishop, 1988). The flux of particulate BaSO<sub>4</sub> to the seafloor is correlated with the flux of exported organic matter (e.g., Dymond et al., 1992; Eagle et al., 2003; Serno et al., 2014; Hayes et al., 2021). This correlation means that the accumulation rate of sedimentary BaSO<sub>4</sub>—or its main constituent, Ba—can be used to trace patterns of past organic matter export on timescales ranging from millenia to millions of years (e.g., Bains et al., 2000; Paytan & Griffith, 2007; Schmitz, 1987; Schroeder et al., 1997).



**Figure 1. Distribution of barium in seawater. A.** Property–property plot showing the 4,345 co-located, core-feature complete dissolved data used in ML model training (Sect. 2). Sample locations shown in Figure 2. Dashed line shows best-fit linear regression through these data, whereby [Ba] =  $0.54 \cdot [Si] + 39.3$ . Panels **B., C., D.,** and **E.** show average Pacific Ocean dissolved depth profiles of [Si], [Ba], Ba\*, and  $\Omega_{\text{barite}}$ , respectively. Solid line denotes the arithmetic mean and the shaded region encompasses one standard deviation either side of the mean. Dashed line indicates Ba\* = 0 (D) and  $\Omega_{\text{barite}} = 1$  (E).

While the Ba-based proxies are valuable, their applications are potentially limited by insufficient knowledge of the distribution of [Ba]. For example, there is significant vertical and spatial variability in the Ba–Si relationship (Sect. 3.3.; Fig.), 1), which we quantify using Ba\* (<u>barium-star;</u> e.g., Horner et al., 2015):

$$Ba^* = [Ba]_{in \ situ} - [Ba]_{predicted}$$
[Eq. 1]

where [Ba]<sub>predicted</sub> is based on the Ba–Si linear regression (Fig. 1):

$$[Ba]_{predicted} = 0.54 \cdot [Si]_{in \ situ} + 39.3$$

$$[Eq. 2]$$

Here,  $[Si]_{in \ situ}$  has units of  $\mu$ mol kg<sup>-1</sup> and  $[Ba]_{predicted}$  nmol kg<sup>-1</sup>; therefore, Ba\* also has units of nmol kg<sup>-1</sup>. The vertical profile of Ba\* is rarely conservative (Fig. <u>41D</u>) and these variations could introduce uncertainty in the reconstruction of [Si] using Ba.

The relationship between sedimentary BaSO<sub>4</sub> accumulation rates and productivity also contains a significant degree of scatter (e.g., Serno et al., 2014; Hayes et al., 2021). Some of this scatter may relate to variability in BaSO<sub>4</sub> preservation, which is at least partially sensitive to ambient saturation

state,  $\Omega_{\text{barite}}$  (e.g., Schenau et al., 2001; <u>Singh et al., 2020;</u> Fig. 1). The saturation state of a parcel of water with respect to BaSO<sub>4</sub> is defined as:

$$\Omega_{\text{barite}} = Q / K_{\text{sp}}$$
[Eq. 3]

where Q is the Ba and sulfate ion product and  $K_{sp}$  is the *in situ* BaSO<sub>4</sub> solubility product. Discerning the importance of  $\Omega_{barite}$  on BaSO<sub>4</sub> preservation has hitherto been challenging owing to the sparsity of *in situ* [Ba] measurements. Accurately determining the global distribution of [Ba] would be valuable for geochemists and oceanographers, and would enable a more thorough investigation of the effects of preservation on BaSO<sub>4</sub> fluxes and refinement of the Ba–Si nutrient proxy.

A powerful way of interrogating oceanic element distributions is through modeling. Broadly, there are two modeling approaches relevant for simulating [Ba]: mechanistic (i.e., theory driven) and statistical modeling (i.e., data driven; e.g., Glover et al., 2011). Mechanistic In mechanistic or process-based modeling-is generally viewed as the gold-standard approach;, model outputs are derived from sets of underlying equations that are based on fundamental theory. As such, mechanistic model outputs can be interrogated to obtain understanding of processes and their sensitivities. However, creating a mechanistic model of the marine Ba cycle requires embedding a biogeochemical model of BaSO<sub>4</sub> cycling within a computationally expensive global circulation model. Although the computational cost associated with building mechanistic models has been reduced by the development of ocean circulation inverse models (e.g., DeVries, 2014; John et al., 2020), this approach still requires detailed parametrizations of the marine Ba cycle, which do not currently exist. In contrast, statistical models are based on extracting patterns from existing data and using those relationships to make predictions. Statistical models encompass a wide variety of approaches ranging from regression analysis to machine learning (ML). Of particular interest to our study are ML models, which can make predictions without any explicit parameterizations of causal relationships. Machine learning models are computationally efficient and can be highly accurate, though they offer limited interpretability. Machine learning is increasingly being used to solve problems in Earth and environmental sciences, including simulating the dissolved distribution of tracers in the sea (e.g., for cadmium, Roshan & DeVries, 2021; copper, Roshan et al., 2020; iodine, Sherwen et al. 2019; nitrogen isotopes of nitrate, Rafter et al., 20192019; and zinc, Roshan et al., 2018).

The goal of this study is to obtain an accurate simulation of [Ba], which ML makes possible even in the absence of a process-level understanding of the marine Ba cycle. We tested thousands of ML models that were trained using quality-controlled GEOTRACES data from the Arctic, Atlantic, Pacific, and Southern Oceans, supplemented by Argo, satellite chlorophyll, and bathymetry data products (Sect. 2.). Models were tested for their accuracy by simulating [Ba] in the Indian Ocean and comparing predictions against observations made between 1977–2013. Importantly,Since no Indian Ocean data were seen by any of the models during training (Sect. 2.). From this, we are able to identify models with high generalization performance (Sect. 2.). theWe then identify an optimal set of predictor variables that results in the most accurate estimates of [Ba],<sub>a</sub> calculate model uncertainties, and simulate [Ba], Ba\*, and  $\Omega_{\text{barite}}$  on a global basis (Sect. 5.). This result will be valuable for researchers interested in marine Ba cycling, and demonstrates the utility of ML to tackle problems in marine biogeochemistry.

# 2. Training and testing data

Machine learning algorithms are adept at making accurate predictions of a target variable by identifying relationships between variables within large data sets. However, making accurate predictions first requires that a ML algorithm is trained on existing observations of that variable alongside a number of other parameters. These other parameters, hereafter termed features, are an important part of model training; features should encode information that may help the ML algorithm predict [Ba], otherwise their inclusion may diminish model performance. Features should also be well characterized in the global ocean, which allows ML models to make predictions in regions beyond the initial training dataset. We selected 12 model features by considering the tradeoff between feature availability and presumed predictive power (Table 1). While testing more features may have resulted in a more accurate final model, we found that many observations of [Ba] did not have corresponding data for <u>severalmultiple</u> features. Thus; thus, including more features would have meant fewer training data. In <u>subsequent sectionsMoreover</u>, we find that only 4–7 including more than nine features are needed to accurately predict [Ba]-can actually diminish model performance. As such, we did not evaluate the predictive power of other <u>predictorsfeatures</u> beyond the initial feature set.-12 initially selected.

 Table 1. List of oceanographic parameters chosenselected as model features. The features tested

 were selected based on their presumed predictive power and geospatial coverage.

#	Parameter Name	Abbreviation	Units	Coverage*
1	Latitude	Lat.	degrees north (°N)	_
2	Longitude	Long.	degrees east (°E)	_
3	Sample collection depth	Z	meters (m)	-
4	Temperature	Т	degrees Celsius (°C)	97.44%
5	Salinity	S	unitless, but often written in 'units' of PSU or PSS	97.44%
6	Dissolved oxygen	[O <sub>2</sub> ]	µmol kg <sup>-1</sup>	97.44%
7	Dissolved nitrate	[NO <sub>3</sub> -]	µmol kg <sup>-1</sup>	97.44%
8	Dissolved phosphate	[ <del>PO</del> 4 <sup>3-]</sup> <u>PO4</u> ]	µmol kg <sup>-1</sup>	97.44%
9	Dissolved silicon (as silicic acid)	[Si <del>(OH)</del> 4]	µmol kg <sup>-1</sup>	97.44%
10	Maximum monthly mean mixed-layer depth	MLD	meters (m)	88.20%
11	Mean average annual surface chlorophyll	Chl <u>.</u> a	mg m <sup>-3</sup>	93.95%
12	Bathymetry	Bathy.	meters (m)	100%

\*Coverage values represent the percentage of data points within the World Ocean Atlas 2018 grid that have available data for a given parameter. Latitude, longitude, and depth have 100 % coverage as these features define the grid itself.

The 12 features used to predict [Ba] and their associated data sources are summarized in Table 1 and described below. The first three features (latitude, longitude, depth) record geospatial information that defines the location of an observation in three-dimensional space. To avoid numerical discontinuities, latitude and longitude were introduced into the model as a hyperparameter consisting of the cosine and sine of their respective values (in radians). Data for features 1–3 were included in the sample metadata. Features 4–9 encode physical (temperature, salinity) and chemical (oxygen, nutrients) information that is routinely measured alongside [Ba]. These data were generally available for the same bottle as the [Ba] measurements; however, when

that was not the case, nutrient data were taken from the corresponding location during a separate cast, or, in the case of oxygen, from linearly interpolated sensor data. The final three features are independent of depth, meaning that all samples within a given vertical profile exhibit the same value for MLD (mixed-layer depth), sea-surface chlorophyll a, and bathymetry. Features 10–12 were drawn from several data sources. A climatology of MLD (feature 10) was compiled using the Argo database (Holte et al., 2017). We selected maximum monthly mean MLD as the feature of interest, as this appears to be the spatiotemporal scale most relevant for influencing [Ba] distributions (Bates et al., 2017). Feature 11 represents a blended SeaWiFS and MODIS climatology of chlorophyll a that was obtained from the Copernicus Marine Environment Monitoring Service (CMEMS, 2021). We calculated the mean annual chlorophyll a for each grid cell in the data product and log transformed the data to reduce parameter weighting (e.g., Rafter et al., 2019). Data for MLD and chlorophyll a were extracted at the location of [Ba] observations using nearest-neighbor interpolation and their values logged in the master record. Bathymetric information (feature 12) was extracted from one of two sources. Our preferred source was the sample metadata, which generally included a value for bathymetry. For samples lacking bathymetric information, we used nearest-neighbor interpolation to extract a value from the ETOPO5 Global Relief Model (National Geophysical Data Center, 1993). Occasionally, the ETOPO5-extracted bathymetry was shallower than the deepest observation of [Ba] in a given vertical profile. In such cases, the bathymetry logged in the master record was set to 1.01 times the depth of the deepest observation in that profile.

The [Ba] data from the Indian Ocean were collected from a multitude of several, primarily pre-GEOTRACES sources (Table 2). As such, these data were generally incomplete for the 12 features used to train the ML models. Rather than using a mixture of *in situ* and interpolated data, we decided to interpolate all Indian Ocean data for parameters 4–12. Data for parameters 4–9 were linearly interpolated from the nearest vertical profile in the World Ocean Atlas 2018 (WOA; Boyer et al., 2018; García et al., 2018a; 2018b; Locarnini et al., 2018; Zweng et al., 2018) and values for MLD and chlorophyll *a* were extracted from the aforementioned data products using nearestneighbor interpolation. Bathymetric information was obtained from either the WOA or *ETOPO5*. For the vast majority of most samples, bathymetry was taken as the arithmetic mean of the maximum depth of the nearest vertical profile in the WOA and the depth at the standard level below. For example, if the maximum depth at a station was 950 m, the bathymetry was recorded as 975 m, which is the mean of levels 46 (950 m) and 47 (1,000 m). For profiles with a maximum depth of 5,500 m—level 102, the lowest in the WOA—bathymetry was recorded as either 5,550 m or the nearest-neighbor interpolated value from *ETOPO5*, whichever was deeper.

Purpose	Region	Expedition ID	Data source	Data Originators (if unpublished)		
	South Atlantic (Meridional)	GA02	GEOTRACES IDP 2017 (Schlitzer et al., 2018)	Jose M. Godoy		
	North Atlantic (Zonal)	GA03	Rahman et al., 2022			
	South Atlantic (Zonal)	GA10	Horner et al., 2015; Bates et al., 2017; Hsieh & Henderson, 2017; Bridgestock et al., 2018			
Model training	Southern Ocean (Meridional)	GIPY04	GEOTRACES IDP 2017 (Schlitzer et al., 2018)	Frank Dehairs		
	Southern Ocean (Zonal)	GIPY05	Hoppema et al., 2010			
	Arctic	GIPY11	Roeske et al., 2012			
		GN01	Whitmore et al., 2022			
	Pacific (Meridional)	GP15	GEOTRACES IDP 2021 (GEOTRACES IDP Group, 2021)	Laura Whitmore, Melissa Gilbert, Emilie Le Roy, Tristan Horner, Alan Shiller		
	Subtropical South Pacific (Zonal)	GP16	Rahman et al., 2022			
		GEOSECS	Craig & Turekian (1980)			
Model	Indian Ocean	INDIGO 1	Jeandel et al. (1996)			
validation		INDIGO 2				
		INDIGO 3				
		SR3	Jacquet et al. (2004)			
		SS259	Singh et al. (2013)			

Table 2. Data sources. Information regarding the source of [Ba] incorporated into the master record.

This data ingestion process resulted in a master record containing 5,502 observations of [Ba] that also contained a corresponding value for all 12 of thecore features listed in (Table 1-). The record was then split into a Pareto partition: the first partition was used for ML model training (4,345 observations, 79 % of data; Fig. 1A) and the second for model testing (1,157 data; 21 %). This partitioning was determined based on the basin from which the sample was collected; data from the Arctic, Atlantic, Pacific, and Southern Oceans were used in model training, whereas the 1,157 [Ba] data from the Indian Ocean were reserved for model testing (Table 2; Fig. 2). This location-based separation of training and testing data was chosen to minimize overfitting, which can occur when the training–testing separation is randomly assigned (see e.g., Rafter et al, 2019).Sect. 3.2.).



**Figure 2. Geographical distribution of the training and testing data**. The 4,345 core-feature complete training data (red; Fig. 1) are from the GEOTRACES 2021 Intermediate Data Product (GEOTRACES IDP Group, 2021); GEOTRACES expedition identifiers are noted next to each section. TestingThe n = 1,157 testing data from the Indian Ocean are color-coded by expedition (see key; n = 1,157); data. Data sources are listed in Table 2.

# 3. Methods

In the following subsections we discuss details of the specific ML algorithm that was used for model development (Sect. 3.1.), explain the model training and testing process (Sect. 3.2.), and describe how a global prediction of [Ba] was obtained and interrogated (Sect. 3.3.).

#### 3.1. Algorithm selection and training

We opted for supervised ML using a Gaussian Process Regression learner, implemented in MATLAB. This particular ML algorithm is non-parametric, kernel-based, and probabilistic<sub>7</sub>, which means that it does not make strong assumptions about the mapping function, can handle nonlinearities, and takes into account the effect of random occurrences when making predictions. Gaussian Process Regression algorithms are widely used in geostatistics, where it is often referred to as 'kriging' (e.g., Cressie, 1993; Rasmussen & Williams, 2006; Glover et al., 2011). This type of algorithm is ideal when working with continuous data that also contains a certain level of noise, such as from measurement uncertainty or oceanographic variation. The basis and kernel function parameters were chosen as constant and exponential, respectively, as this combination was found to produce the most accurate predictions. The MATLAB function, fitrgp, was used for model training. A full list of the parameter selections used in fitrgp is provided in Table S1. All predictors were normalized and standardized to have a mean of zero and a standard deviation of unity. This process placed places all parameters on the same relative range and was intended to diminishreduces scale dependencies.

A significant problem in supervised ML algorithms is overfitting: the tendency to produce highly precise fits to the training data that cannot then be generalized to new domains or environments. We attempted to minimize overfitting by performing cross-validation during model training and during model testing. First, we used holdout cross folding during model training. Data were randomly split into two folds, one containing 80 % of the data for model training and the other 20 % withheld for model validation (i.e., the holdout fold). This holdout process was intended to eliminate models that could only generate arbitrary fits to specific subsets of the training data. In the second stage of cross-validation, we evaluated the performance of trained models by comparing predictions against a set of withheld [Ba] observations from the Indian Ocean. None of

the Indian Ocean data were seen by the models during training. This withholding was intended to help identify models that were generalizable to new environments and therefore the entire ocean (see Sect. 3.2).

#### 3.2. Model training and testing

The training partition of the master record was used to train 4,095 different machine learning models with the goal of finding a model that could accurately simulate the global distribution of [Ba]. The number of models tested derives from the number of features investigated; each model uses a unique combination of the 12 features in Table 1 and our testing followed a factorial design whereby each feature was either enabled or disabled. This design yields a total of  $2^{12}$  unique feature combinations (i.e., levels<sup>features</sup>). Since); however, since it is not possible to train a model with no features enabled cannot be trained, the final number of unique, trainable, ML models with  $\geq 1$  features wasis  $2^{12}-1=4,095$ . The full experiment list is provided in Section 6. Each of the 4,095 models was trained using the same 4,345 input data and with the same function parameters described above (Sect. 3.1.). Testing every possible feature combination allowed us to select for models with the highest predictive power while minimizing overfitting. in Table S1.

<u>3.2.</u> In the second stage of cross validation, trained models were used to predict [Ba] for the withheld data from the Indian Ocean. Each of the 4,095 trained models were provided with the feature information that that particular model required to simulate [Ba] and the predictions recorded. The accuracy of the models was assessed by comparing ML model predictions against observed [Ba] for the Indian Ocean data (n = 1,157) and calculating the mean absolute deviation (MAD) and mean absolute percentage deviation (MAPD). The MAD is defined as:

# MADAssessing model performance

Model performance—accuracy and generalizability—was assessed during two phases: training and testing. During model training, the 4,345 observations of [Ba] from the Arctic, Atlantic, Pacific, and Southern Oceans were randomly split into two folds: a training fold containing 80 % of the observations, and a holdout fold containing the other 20 %. Model accuracy was assessed by comparing model-predicted [Ba] against observed [Ba] for the 20 % of the data in the holdout

fold. We then performed additional testing to establish model generalizability. A significant problem in supervised ML, and particularly Gaussian Process Regression learning, is overfitting: models may fit the noise in the training data, leading to poor generalization performance (Rasmussen & Williams, 2006). Since our goal was to develop a global model of [Ba] using regional training data, we deemed it especially important to identify generalizable models. Generalizable models were identified through a testing process involving regional cross-validation; each trained model was used to predict [Ba] for the 1,157 samples from the Indian Ocean and model predictions were again compared against observations. Importantly, no [Ba] data from the Indian Ocean were seen by any of the models during training. This process helped to identify models that may have been overfit to the training data and can further be used to calculate generalization errors (Sect. 4.1).

The accuracy of trained models was determined by comparing ML model predictions against withheld data and calculating the mean absolute error (MAE) and mean absolute percentage error (MAPE), defined as:

$$\underline{MAE} = \frac{\sum_{i=1}^{n} |[Ba]_{predicted} - [Ba]_{observed}|}{n} \sum_{i=1}^{n} |[Ba]_{predicted} - [Ba]_{observed}|}{n}$$
[Eq. 4]

and MAPD as:

$$\frac{\text{MAPD}}{\text{MAPE}} = \frac{100 \%}{n} \frac{\sum_{i=1}^{n} \left| \frac{[Ba]_{predicted} - [Ba]_{observed}}{[Ba]_{observed}} \right|}{[Ba]_{observed}} \sum_{i=1}^{n} \left| \frac{[Ba]_{predicted} - [Ba]_{observed}}{[Ba]_{observed}} \right|$$
[Eq. 5]

# respectively, where *n* is the sample size.

Models with lower accuracy exhibit higher MAD and MAPDerrors, whereas models with high accuracy will-have lower MAD and MAPD. For reference, the Ba–Si linear regression predicts Indian Ocean [Ba] with a MAD and MAPDerrors. We calculated MAE and MAPE for every possible feature combination, which enables quantification of 6.8 nmol kg<sup>-1</sup> and 9.7 %, respectively (Fig. 3). These values can be considered as benchmarks for the ML models. how specific features affect model performance. Likewise, we calculated errors for each model on predictions made during training (i.e., for the holdout fold) and during model testing (i.e., during regional cross-validation; Fig. 3). This information is used to quantify generalization performance;

low errors for both training and testing indicate models that are both accurate and generalizable, whereas models with low training errors and high testing errors might indicate models -overfit to the training data.

# 3.3. Global predictions

A select number of models with low MADMAE and MAPDMAPE were used to simulate [Ba] on a global basis. The process by which we selected these models is described in Section 5.1. Global simulations were performed on the same grid as the WOA, which was also used as the data source for features 1–9 (Boyer et al., 2018). The WOA is a  $1^{\circ}\times1^{\circ}$  resolution data product with around 41,000 stations that contain up to 102 depth levels spanning 0–5,500 m in 5, 25, 50, or 100 m increments. Data for features 10–12 (MLD, chlorophyll *a*, and bathymetry) were also resampled to the WOA grid using the same sources and interpolation methods as described for the Indian Ocean testing data in Section 2. Model outputs were visualized using Ocean Data View software (ODV; Figs. 4–75–8; Schlitzer, 2023).

A selection of the most accurate models of [Ba] were then used to simulate Ba\* and  $\Omega_{\text{barrite}}$ . The ealeulationA selection of the most accurate models of [Ba] were then used to simulate Ba\* and  $\Omega_{\text{barrite}}$ . Star tracers, such as Ba\*, are valuable for illustrating processes that influence the cycling of elements in the ocean. First defined for N–P decoupling (N\*; Gruber & Sarmiento, 1997) star tracers show variations whenever there are differences in the sources and sinks of the two elements being compared. If there are no differences in sources and sinks, the tracer will show conservative behavior because both elements share the same circulation. Barium-star is based on Ba–Si decoupling and was first defined by Horner et al. (2015). The definition of Ba\* is shown in Equations 1 and 2. The coefficients in Equation 2 are based on data from the GEOTRACES 2021 Intermediate Data Product and specifically the subset of these data shown in Figure 1. These coefficients differ from previous formulations of Ba\* that were based primarily on [Ba] and [Si] data from the Southern and Atlantic Oceans (e.g., Horner et al., 2015; Bates et al., 2017). CalculationThe global distribution of ML-model derived Ba\* used values of [was determined by calculating [Ba]<sub>predicted</sub> (Eq. 2) using [Si]<sub>in situ</sub> from the WOA 2018 (García et al., 2018b<del>) and</del>). The

# values of [Ba]<sub>in situ</sub> was taken from the ML model output- and [Ba]<sub>predicted</sub> was subtracted from this to yield Ba\* (Eq. 1).

Values of  $\Omega_{\text{barite}}$  were computed using the method described by Rushdi et al. (2000), summarized in Equation 3. In The numerator, Q, represents the in situ Ba and sulfate ion product and, in this formulation, sulfate is assumed to be conservative with respect to salinity and thus this method cannot be used to predict  $\Omega_{\text{barite}}$  in restricted basins, such as the Black Sea or Caspian Sea. As with depends only on [Ba] and [SO<sub>4</sub><sup>2-</sup>] molality. The denominator,  $K_{sp}$ , depends on T, S, and z (i.e., pressure) and is calculated in two steps: in situ T and S are used to calculate the stoichiometric solubility product and then this value is modified by calculating the effect of pressure on partial molal volume and compressibility, which are functions of T and z. As with the calculation of Ba\*, values of [Ba]in situ were obtained from ML models and co-located data for T, S, and pressure dataz were extracted from the WOA (Locarnini et al., 2018; Zweng et al., 2018). Sulfate concentrations were assumed to be conservative with respect to S using  $[SO_4^{2-}] = 29.26$  mmol kg<sup>-1</sup> when salinity = 35 PSU. This latter assumption likely breaks down in certain environments (e.g., where  $[SO_4^{2-}]$ ] reduction occurs); as such, our model is not used to predict  $\Omega_{\text{barite}}$  in restricted basins, such as the Black Sea or Caspian Sea. Given that our estimates of  $\Omega_{\text{barite}}$  exhibit a MAE of 0.08 (Appendix), we believe that values of  $\Omega_{\text{barite}}$  between 0.92 and 1.08 are indicative of 'perfect' saturation with respect to BaSO<sub>4</sub>.

Output from the most accurate ML models was then used to calculate mean [Ba] and  $\Omega_{\text{barite}}$  for each basin, for a series of prescribed depth bins, and for the global ocean. This calculation was performed by weighting each cell in the model output by its volume, which ensures a fair comparison between any two points in the model output. We then subdivided the global ocean into five sub-basins: Arctic, Atlantic, Indian, Pacific, and Southern. Basin boundaries were defined as per Eakins & Sharman (2010), though we merged the Mediterranean and Baltic Seas into the Atlantic and considered the South China Sea as part of the Pacific Ocean. Neither [Ba] nor  $\Omega_{\text{barite}}$ were simulated in the Black or Caspian Seas and thus these regions are not included in the global mean calculations.

# 4. Results

#### 4.1. Quantifying Factors affecting model accuracy

Here we examine model accuracy and assess the role of different features in settinghow model performance. Accuracy was assessed using the mean absolute deviation (MAD; Eq. 3), which is a measure of the correspondence between predicted and observed [Ba] for the n = 1,157 data from the Indian Ocean. This correspondence is illustrated for [Ba] predicted using WOA-interpolated [Si] and the Ba-Si linear regression (Fig. 3A) and for ML model #3336 (Fig. 3B; Sect. 5.1.). The calculation of MAD was repeated for all 4,095 trained models and the results are summarized in Figure 3C. Of these models, 1,687 (41%) achieve a superior MAD in the Indian Ocean compared to the Ba-Si linear regression benchmark of 6.8 nmol kg<sup>-1</sup>. In general, ML models with fewer features tend to exhibit higher MAD than models with many features. However, adding more features to a model can also degrade its performance. Binnedinfluenced by the number of features, the MAD of the median model decreases from 15.8 to 7.1 and nature of features included during training. We consider model performance in terms of accuracy and generalizability, which we guantify using MAE (Eq. 4). We first explore how the number of features influences model performance (Fig. 3). Here we see that increasing the number of features generally improves the accuracy of trained models; however, the response differs depending on whether accuracy is calculated based on comparison to the holdout fold (i.e., during model training) or to the withheld Indian Ocean data (i.e., during model testing). When considering only the holdout fold, trained models predict [Ba] with a high level of accuracy-the mean, median, and most-accurate trained models achieve a MAE of 2.4, 1.7, and 1.3 nmol kg<sup>-1</sup>, respectively. Similarly, increasing the number of features almost always improves model accuracy; the MAE of the most accurate model for a given number of features decreases from 6.5 to 1.3 nmol kg<sup>-1</sup> as the number of features is increased from one to five. Beyond five features, the median-model MADnine, at which point MAE plateaus; the median MAD of the 2,510 models with  $\geq 6$  features is 7.8 nmol kg between 1.4–1.5 nmol kg<sup>-1</sup> for models with 10–12 features (Fig. 3A).



**Figure 3. Effect of feature addition on ML model accuracy.** Accuracy was quantified for each of the 4,095 trained models and quantified here using MAE (note log scale, which differs between panels). The accuracy of trained models is shown for random holdout cross-validation during training (top) and for regional cross-validation during testing (bottom). Square indicates the performance of our favored predictor model, #3080 (see Fig. 4, Sect. 5.1). The accuracy of the Ba–Si linear regression benchmark is shown as a dashed line in the lower panel (MAE = 6.8 nmol kg<sup>-1</sup>). To illustrate data density, points have been randomly positioned within their respective bin and plotted with 80 % transparency.

Moving to the regional cross-validation, the overall performance of models is lower; the same 4,095 trained models achieve a mean, median, and most-accurate MAE for the Indian Ocean dataset of 8.8, 7.9, and 4.0 nmol kg<sup>-1</sup>, respectively. For comparison, if [Ba] was estimated for these

same 1,157 Indian Ocean samples using the linear [Ba]–[Si] relationship (Fig. 1) and ambient [Si] as the only predictor, this linear model would achieve a MAE of 6.8 nmol kg<sup>-1</sup>. Thus, there are 1,687 ML models that achieve a superior accuracy to existing methods for estimating [Ba], offering an improvement of as much as 41 % (Fig. 4). However, regional cross-validation also shows that the addition of more features may, in fact, degrade model performance. The MAE of the most accurate model for a given number of features decreases from 6.6 to 4.0 nmol kg<sup>-1</sup> as the number of features is increased from one to eight. As the number of features is increased from nine to twelve9–12, the MADMAE of the most-accurate ML model within each bin monotonicallymodels increases monotonically from 4.1 to 7.2 nmol kg<sup>-1</sup> (Fig. 3C). Thus, the number of features necessary1 nmol kg<sup>-1</sup>. The overall lower performance of trained models during regional cross validation—and the observation that many of the feature-rich models perform worse than models with fewer features—is indicative of certain models being over-fit to the training data. Together, these observations suggest that the optimum number of features necessary predict [Ba] in the Indian Ocean appears to beis between five and eight.

We then quantified the importance of different features to model performance through a feature addition analysis (Fig. 3D). For example, model #3352 contains five features: z, S,  $[O_2]$ ,  $[PO_4]$ , and  $[NO_3]$  and achieves a MAD of 4.6 nmol kg<sup>-1</sup>. Adding T to this model increases the number of features to six and <u>nine</u>.



**Figure 4. Comparison of existing and ML methods to estimate [Ba] in seawater.** Left panel shows the performance benchmark: predicted [Ba] for the Indian Ocean testing data using the [Ba]–[Si] linear regression and ambient [Si] as the sole predictor. Right panel shows predicted [Ba] using ML model 3080, which improves on existing methods by more than 37 %. Perfect correspondence between predictions and observations is indicated b the dashed line marked '1:1.'Data locations and sources are shown in Fig. 2 and Table 2, respectively; *n* refers to the number of testing data for each campaign. reduces Mean Absolute Error (MAE; Eq. 4) and Mean Absolute Percentage Error (MAPE; Eq. 5) are noted for both models.

We also evaluated the MAD to 4.4 nmol kg<sup>-1</sup> (Fig. 3B). Since wenature of the predictors used ato estimate [Ba]. The full factorial experiment design, we were able enables us to perform analogous pairwise comparisons between the 2<sup>++</sup>all models that contained a certain feature, such as *T*, and the 2<sup>++</sup>all of those that did not- (Sect. 3.1). We quantified the effect of adding a feature by comparing the absolute and percentage change in MADMAE relative to the mean MADMAE of the two models. The likelihood that the inclusion of a given feature affected the MAD of the models sets of models. This comparison was then quantified using a two tailed, paired *t* test. Lower *p* values indicate performed three times: for all 4,095 models based on the holdout crossfolded training data, for all models using the regionally cross-validated testing data, and again for the testing data, but only considering those 1,687 models that achieved a higher likelihood that superior accuracy compared to the [Ba]–[Si] linear regression model (Table 3).

<u>Table 3. Feature</u> addition <u>analysis</u>. Effect of <u>aeach</u> feature <u>significantly changed the MAD</u>. <u>on model</u> performance for Training and Testing datasets. Model performance is quantified using MAE, thus all columns have units of nmol kg<sup>-1</sup> unless otherwise shown. The Testing analysis is further subdivided into a comparison of all models and 'good' models, meaning those that achieved superior accuracy than the Ba–Si linear regression (Fig. 1).

	Training		Testing							
	All models ( <i>n</i> = 4,095)		All models ( <i>n</i> = 4,095)		Good models ( <i>n</i> = 1,687)					
Feature	Mean MAE of models with feature	Mean MAE of models without feature	Relative change in MAE	Mean MAE of models with feature	Mean MAE of models without feature	Relative change in MAE	Mean MAE of models with feature	Mean MAE of models without feature	Relative change in MAE	Share of models with feature
[Si]	1.71	3.03	-56%	7.08	10.6	-39%	5.06	5.50	-8.3%	63%
z	1.83	2.90	-45%	7.94	9.70	-20%	5.05	5.44	-7.4%	55%
[O <sub>2</sub> ]	2.03	2.71	-29%	8.25	9.39	-13%	5.14	5.33	-3.8%	54%
Τ	1.78	2.96	-50%	7.61	10.0	-27%	5.17	5.31	-2.8%	59%
[NO <sub>3</sub> ]	2.09	2.65	-24%	8.27	9.36	-12%	5.16	5.30	-2.7%	53%
[PO₄]	2.11	2.63	-22%	8.24	9.40	-13%	5.17	5.30	-2.4%	53%
S	2.02	2.72	-29%	8.67	8.97	-3.5%	5.23	5.23	0.0%	53%
Bathy.	2.30	2.44	-6.1%	8.55	9.08	-6.0%	5.23	5.22	0.2%	51%
Chl.	2.25	2.48	-10%	8.67	8.97	-3.5%	5.24	5.22	0.4%	50%
MLD	2.31	2.43	-4.8%	8.69	8.95	-3.0%	5.24	5.21	0.5%	50%
Lat.	2.16	2.58	-18%	8.13	9.51	-16%	5.32	5.11	4.0%	54%
Long.	2.17	2.57	-17%	11.4	6.24	58%	6.45	5.19	22%	3%

This analysis reveals that yields three main results. When considering only the holdout cross-folded training data, the addition of any one of 11 features will, on average, improve an ML model. Silicate ( 40 %), *T* ( 27 %), and *z* ( 20 %) of the 12 features improves model performance by between –4.8 and –56 %. Excepting longitude, similar across-the-board improvements were observed when considering only the testing data, though the improvements for most features were more modest (between –3.0 and –39 %). If considering only the 'good' models, six features improved the models the most and *S*, chlorophyllmodel performance by –2.4 and –8.3 % ([PO<sub>4</sub>], [NO<sub>3</sub>], *T*, [O<sub>2</sub>], *z*, and [Si]), five degraded model performance by +0.2 to +22 % (bathy., Chl. a, and MLD the least (all –3 %). Latitude, [PO<sub>4</sub>], [O<sub>2</sub>], [NO<sub>3</sub>], and bathymetry improved the models by –16 % to –6 %. Longitude was the only feature found to degrade model performance, with a mean change in MAD of +59 %. The largest *p* value associated with these comparisons was 5E–25, indicating that these relationships were highly, lat., and long.), and salinity had no significant (Fig. 3Deffect (Table 3).

Overall, our results indicate that between six and nine features will result in an accurate and generalizable ML model of [Ba], and that  $[PO_4]$ ,  $[NO_3]$ , *T*,  $[O_2]$ , *z*, [Si], and possibly *S*, are likely to be included as predictors in such a model.

# 4.2. Model outputs

Almost 1,700 models achieved superior accuracy compared to the Ba–Si linear regression benchmark of 6.8 nmol kg<sup>-1</sup>. We winnow this list to a single model, #33363080, in the next section. We henceforth refer to model #33363080 as our favored predictor model, which achieves a MADMAE of 4.43 nmol kg<sup>-1</sup> using *z*, *T*, *S*, [O<sub>2</sub>], [PO<sub>4</sub>], [NO<sub>3</sub>], and [NO<sub>3</sub>Si] as predictors (Fig. 3B4). Model #33363080 is used to simulate [Ba], Ba\*, and  $\Omega_{\text{barite}}$  on a global basis and to calculate whole-ocean averages. Surface plots showing the model outputs for the sea surface, 1,000 m, 2,000 m, and 4,000 m are shown in Figures 4, 5, 6, 7, and 78, respectively.



**Figure 45**. **Barium at the sea surface.** Observed [Ba] between 0–50 m (**A**); Model 33363080 [Ba] (**B**), Ba\* (**C**), and  $\Omega_{\text{barite}}$  (**D**). The dashed line in Panel D indicates the BaSO<sub>4</sub> saturation horizon (i.e.,  $\Omega_{\text{barite}} = 1.0$ ). Panels A and B use the *roma* color map, whereas Panels C and D use *vik* and *cork*, respectively (Crameri, 2018). Color palettes and parameter ranges are the same for the respective panels in Figure 5–76–8.



<u>Figure 6. Barium at 1,000 m.</u> Observed [Ba] (A); Model  $\frac{33363080}{1}$  [Ba] (B), Ba\* (C), and  $\Omega_{\text{barite}}$  (D). The dashed line in Panel D indicates the BaSO<sub>4</sub> saturation horizon.



**Figure 7. Barium at 2,000 m.** Observed [Ba] (**A**); Model 3080 [Ba] (**B**), Ba\* (**C**), and Ω<sub>barite</sub> (**D**). The dashed line in Panel D indicates the BaSO<sub>4</sub> saturation horizon.



**Figure 7. Barium at 4,000 m.** Observed [Ba] (**A**); Model  $\frac{33363080}{33363080}$  [Ba] (**B**), Ba\* (**C**), and  $\Omega_{\text{barite}}$  (**D**). The dashed line in Panel D indicates the BaSO<sub>4</sub> saturation horizon.

Model #33363080 contains 3,305,505302,570 predictions for each of [Ba], Ba\*, and  $\Omega_{\text{barite}}$ . Values (Sect. 6). Assuming that the MAPE and MAE are good estimates of the prediction error, we estimate that modeled [Ba] range from 23.3–158 and Ba\* have uncertainties of 6.0 % and 4.3 nmol kg<sup>-1</sup>, with an unweighted mean of 71 respectively. Uncertainties on  $\Omega_{\text{barite}}$  were estimated by comparison to literature data, which yields a MAE of 0.08. These estimates are discussed in more detail in Section 5.2 and the Appendix.

Modeled [Ba] ranges from 26.2–156.8 nmol kg<sup>-1</sup>- and the data exhibit an unweighted mean of 72.0 nmol kg<sup>-1</sup>. The range of model #3080 predictions is within the range of [Ba] encountered in the 4,345 training data (17.1–159.8 nmol kg<sup>-1</sup>). This is an important consideration when assessing the accuracy of Gaussian Process Regression models, and we provide additional discussion of this point in the Supplement. Based on our formulation of Ba\* (Eqs. 1, 2), Ba\* varies from -102.727.2 to  $+\frac{51.327.9}{1.327.9}$  nmol kg<sup>-1</sup> and possesses an unweighted mean of +2.24 nmol kg<sup>-1</sup>. Values of  $\Omega_{\text{barite}}$ vary from 0.11 to 1.7670 and exhibit an unweighted mean of 0.75. To account for the different volumes represented by each grid cell in the modelWOA grid, we constructed a volume-weighted mean of [Ba] and  $\Omega_{\text{barite}}$  for the ocean as a whole, for each ocean basin, and for a series of prescribed depth bins (Fig. 8). Looking at the ocean as a whole, the probability density function of [Ba] roughly resembles a uniform distribution, with a mean ocean [Ba] of 89 nmol kg<sup>-1</sup> (Fig. 8A9A). Within this mean is considerable spatial and vertical variation. For example, the Arctic Ocean exhibits the lowest volume-weighted mean [Ba] of  $\frac{5554}{554}$  nmol kg<sup>-1</sup>, whereas mean Pacific [Ba] = 106 nmol kg<sup>-1</sup>. Likewise, The Indian Ocean exhibits a similar mean [Ba] exceeding (90 nmol kg<sup>-1</sup>) to the mean of the global ocean. Shallower than 1,000 m, [Ba] infrequently exceeds 100 nmol kg<sup>-1</sup>-rarely occurs above 1,000 m and values, whereas concentrations <45 nmol kg<sup>-1</sup> are virtually absentrare below 1,000 m (Fig. 8B9B).

The probability density function of volume-weighted  $\Omega_{\text{barite}}$  is <u>closermore similar</u> to a normal distribution, <u>possessingalbeit with a slight negative skew. Volume-weighted mean value of oceanic</u>  $\Omega_{\text{barite}}$  is 0.82. The Arctic, Atlantic, and Indian Oceans are, on average, undersaturated with respect to BaSO<sub>4</sub>, all exhibiting <u>mean  $\Omega_{\text{barite}} \leq 0.8182$ </u>. In contrast, the Pacific Ocean is close to and

Southern Oceans are within uncertainty of saturation-(, with mean  $\Omega_{\text{barite}} = \text{of } 0.97$ ), and the Southern Ocean slightly exceeds it ( $\Omega_{\text{barite}} = \text{and } 1.04$ ; respectively (Fig. 8C9C). Values of  $\Omega_{\text{barite}}$ <0.25 are only found above 1,0002 are mostly restricted to the upper 250 m, whilst values of  $\Omega_{\text{barite}}$ exceeding 1.455 are exceptionally rare and are, found only in the upper 1,000 m of the Southern Ocean. Lastly,  $\Omega_{\text{barite}}$  tends to increase between the 0–250 m, 250–1,000 m, and 1,000–2,000 m depth bins, increasing from 0.42, to 0.6365, and 0.96, respectively. Average  $\Omega_{\text{barite}}$  in the deepest bin (2,000–5,500 m) is slightly lower, with a mean value of 0.92 (Fig. 8D).9D). Given the accuracy of our model-derived  $\Omega_{\text{barite}}$  predictions (0.08 to 0.10), the ocean between 1,000–5,500 m is within uncertainty of BaSO4 equilibrium.



Figure 89. Stacked, volume-weighted histograms showing the relative frequency distribution of dissolved [Ba] (A, B) and  $\Omega_{\text{barite}}$  (C, D) in the global ocean. The left column shows data grouped by basin, whereas and the right column is shows data grouped by a prescribed depth bin-(key in B). Numbers in each panel display the mean property value for that bin. Dashed line shows the global mean.

# 5. Discussion

#### 5.1. Identification of the optimal predictor model

Our results show that 1,687 of the 4,095 ML models (41 %) produce more accurate predictions than the benchmark. In this case, the benchmark is predicting Indian Ocean [Ba] based on the Ba-Si linear regression and using WOA [Si] as the only predictor (Figs. 1, 3C). However, choosing the optimal feature combination is challenging given the sheer number of skillful ML models. Here, we winnow the list from 1,687 to a single model (#3336) by eliminating models based on the number and information content of various features. First, longitude was found to be the only feature that consistently degraded the performance of trained models (Fig. 3D). We therefore disabled this feature, eliminating 53 models. Having removed longitude, we also decided to eliminate models utilizing latitude; this decision was guided by the principle that a generalizable model should be able to predict [Ba] using only physical and/or chemical predictors, independent of where a sample is located in x y space. This removed a further 853 models, winnowing the list to 781. After longitude, the features offering the least improvement to ML model performance were MLD and Chl. a, which improved trained models by around -3 % (Fig. 3D). Indeed, the median model was degraded by 2 % if Chl. a was included. Eliminating models containing either of MLD (397) or chlorophyll a (194 models) reduced the number of models exceeding the Ba-Si benchmark of 6.8 nmol kg<sup>-1</sup> to 190, of which 122 utilized [Si] as a predictor. While [Si] was amongst the strongest overall predictors of [Ba] (Fig. 3D), incorporating [Si] into a trained model introduces potential circularity into the calculation of Ba\* (see Eqs. 1, 2). Moreover, the four-most accurate surviving models containing [Si], models 3144, 3268, 3716, and 3732, achieve a similar MAD of 4.2 nmol kg<sup>-1</sup> to our ultimately favored model #3336 (4.4 nmol kg<sup>-1</sup>).

Eliminating models containing [Si] reduced the list of models to 68. Of these, 39 were eliminated as they contained  $\leq$ 4 features, noting that our earlier analysis showed that the optimal number of features needed to accurately predict [Ba] in the Indian Ocean was between five and eight (Sect. 4.1.). Models with fewer than four features likely do not contain sufficient information to make accurate predictions in the full range of environments encountered in the Indian Ocean. Likewise, beyond eight features, trained models tended to produce worse fits to observations. We suspect that the reason is overfitting; feature-rich models can be overfit to the training data and are unable to generalize when presented with completely new environments. Choosing a single, optimal model

configuration is challenging given the sheer number of skillful ML models. Below we winnow the list from 4,095 to a single model (#3080). We base our winnowing primarily on the results of the regional cross-validation performed in the Indian Ocean, rather than from the errors determined from random holdout cross folding of the training data. We believe that there are three strong reasons for winnowing in this way. First, Gaussian Process Regression Learners tend to fit the noise in the training data, meaning that the training error is significantly lower than the generalization error (Rasmussen & Williams, 2006). Indeed, trained models showed overall lower performance during testing compared to training, which we believe is evidence of overfitting (Fig. 3, Table 3). Second, a generalizable global model should be able to make predictions in regions where it has not already learned anything about the target variable. Our regional cross-validation approach satisfies this consideration since no Indian Ocean data were seen during model training. Third, the Indian Ocean is an ideal basin for testing as it exhibits the full diversity of features expected to influence [Ba] (riverine inputs, oxygen-minimum zones, coastal upwelling, etc.) and constitutes  $\approx 20$  % of the global ocean volume. Likewise, the Indian Ocean captures most of the range in [Ba] seen elsewhere in the ocean (Fig. 9); this likely reflects the input of Atlantic waters through the Aughulas leakage, transport of old Pacific waters via the Indonesian Throughflow, and northward spreading of mode and intermediate waters from the Southern Ocean. We thus assume that the Indian Ocean testing errors are a good approximation of the generalization error, which we now use to winnow the list of models.

Our results show that 1,687 of the 4,095 ML models (41 %) produce more accurate predictions of [Ba] than the benchmark Ba–Si linear regression using [Si] as the sole predictor (Fig. 3, Table 3). We focus our winnowing on these 1,687 models as they are superior to existing methods for estimating [Ba] in seawater. Focusing only on these 'good' models reveals significant differences in the information content of the 12 features tested. For example, the inclusion of spatial information in the form of latitude and longitude significantly degrades mean model performance by between +4.0 and +22 %, respectively. While bathymetry, chlorophyll *a*, and mixed-layer depth exhibited only minor influences, they were nonetheless deleterious to mean model performance by between +0.2 to +0.5 % (Table 3). Only [PO<sub>4</sub>], [NO<sub>3</sub>], *T*, [O<sub>2</sub>], *z*, and [Si] consistently improved the mean ML model, which corresponds to model #3112 (testing MAE of 4.3 nmol kg<sup>-1</sup>). However, visual inspection of model #3112 output reveals that it does not reproduce expected near-shore surface plumes of elevated [Ba] close to certain major rivers (see Supplement). Though

volumetrically minor, riverine inputs are a geochemically important component of the marine Ba cycle, and the existence of nearshore Ba plumes underpins a major proxy application of Ba. Nearshore riverine influence is easily discerned by low *S*; we thus explored output from model #3080, which is identical to model #3112, but includes *S* as a seventh feature during training. Models #3080 and #3112 exhibit identical statistical performance for the testing data (MAE = 4.3 nmol kg<sup>-1</sup>; Fig. S1) and make similar predictions for mean marine [Ba] and  $\Omega_{\text{barite}}$  (89 nmol kg<sup>-1</sup> and 0.82, respectively; see Supplement). The similar performance of the two models is consistent with *S* exerting a near-negligible impact on overall model performance (Table 3). Despite this small effect, model #3080 is better able to reproduce riverine [Ba] plumes compared to model #3112 (see Supplement). We therefore consider model #3080 to be our best estimate of marine [Ba]. Model #3080 achieves a MAPE of 6.0 %, which represents a 39 % improvement over existing methods to estimate [Ba] (Fig. 4). We henceforth consider model #3080 as our optimal predictor model, which we use to simulate [Ba], Ba\*, and  $\Omega_{\text{barite}}$  in Figures 5–9.

Thus, this winnowing process reduced the number of candidate ML models to 29. All of these models exhibited superior accuracy compared to the Ba–Si linear correlation (MAD = 6.8 nmol kg<sup>-1</sup>); were not trained using longitude, latitude, MLD, chlorophyll *a*, or [Si]; and possessed between five and seven features. The most accurate amongst these 29 models, #3336, utilizes six features *z*, *T*, *S*, [O<sub>2</sub>], [PO<sub>4</sub>], and [NO<sub>3</sub>] – and achieves a MAD and MAPD in the Indian Ocean of 4.4 nmol kg<sup>-1</sup> and 6.3 %, respectively (Fig. 3B). This level of accuracy represents at least a 35 % improvement compared to predicting [Ba] in seawater using the Ba–Si linear regression. We consider model #3336 as the optimal configuration for predicting [Ba] in this study, and use this model to simulate [Ba], Ba\*, and  $\Omega_{\text{barite}}$  in Figure 4–8.

#### 5.2. Model validation

We now explore the validity of model #33363080 in terms of its oceanographic consistency, the sources of uncertainty that affect its accuracy, and potential limitations of the model output. We find that model #33363080 reproduces the major known features of the marine [Ba] distribution and makes testable predictions for regions that are yet to be sampled.

#### 5.2.1. Visual inspection of model output

Visual inspection of model output is an important component of data analysis considering the limits of statistical tests (see e.g., Anscombe, 1973). Models may produce statistically satisfactory fits to the testing data, but the oceanic realism of the output is also important to consider. Modeled [Ba] should display patterns consistent with related oceanographic properties and exhibit smooth vertical and spatial variations (Boyle & Edmond, 1975). Predicted [Ba] from model #33363080 does indeed show smooth and systematic spatial and vertical variations that also resembles sparse observations (Figs. 3-74-8).

There are, however, several sharp gradients in modeled [Ba], particularly at the sea surface (Fig. 3). These variations generally show an increase Model #3080 also shows systematic increases in [Ba] close to land and, especially near the mouths of major rivers- (Fig. 4). This is reassuring given that elevated sea-surface [Ba] close to rivers is both widely reported and is one of the major proxy applications of Ba: reconstructing spatiotemporal patterns of terrestrial runoff by measuring the Ba:Ca ratio of carbonates (e.g., Sinclair & McCulloch, 2004; LaVigne et al., 2016). Model #3336For example, model #3080 correctly identifies elevated [Ba] near the Ganges–Brahmaputra (Singh et al., 2013) and), Río de la Plata outflows (GEOTRACES IDP Group, 2021), and Yangtze outflows (Cao et al., 2021). Model #33363080 also predicts elevated sea-surface [Ba] in the Gulf of Guinea (where several rivers discharge, including the Niger River; the Eastern Tropical Atlantic associated with the Congo River (Edmond et al., 1978; Zhang et al., 2023); and Volta Rivers) as well as in the Gulf of St. Lawrence (St. Lawrence River), though; see Supplement for additional details and figures). Except for the Congo River, these latter-predictions of elevated near-shore [Ba] await corroboration. Interestingly, model #33363080 does not predict elevated surface-[Ba] at all major outflowsriver mouths; neither the Mississippi nor Amazon Rivers are associated with significant increases in sea-surface [Ba].] (see Supplement). The reasons for the lack of elevated **Ba**] at near the outflow of these rivers two rivers is less clear. It is possible that the model is simply inaccurate in these regions, though we have no particular reason to believe that this is the case. <u>Alternatively, it may reflect seasonal variations in Ba release that are not captured by our mean</u> annual model (e.g., Joung & Shiller, 2014), or it may). It could also indicate that these particular rivers are not major *net* sources of Ba to the surface ocean, which might be the case if dissolved

Ba is being retained in the catchment (e.g., Charbonnier et al., 2020) or estuary (e.g., Coffey et al., 1997).

Overall, model #33363080 makes accurate, oceanographically consistent predictions of [Ba] in the Indian Ocean using input data from the WOA. Model #33363080 also makes a number of testable predictions of [Ba] in regions lacking direct observations. Given that these predictions were made using the same model and the same WOA inputs, we believe that it is reasonable to assume that model #33363080 output is an accurate representation of mean annual global [Ba].

#### 5.2.2. Quantifying uncertainties

We now describe and, where possible, quantify two possible sources of uncertainty to our ML model output. Before doing so, we describe how uncertainty is quantified as well as the uncertainty of existing approaches. Certain ML models, such as Gaussian Process Regression, offer low interpretability, meaning it is not possible to assess uncertainty using a conventional error propagation. Thus, all model uncertainties are assessed *post hoc*, by comparing predictions against observations. Our preferred metrics are MADMAE and MAPD mean absolute deviation and mean absolute percentage deviation, defined in EquationsMAPE (Eqs. 4-and, 5, respectively.). Existing approaches for estimating [Ba] result in a wide range of uncertainties. At the low end, the uncertainty associated with measuring [Ba] in seawater represents a fundamental limit to the accuracy of any model. A number of analysts report relative [Ba] measurement uncertainties in the range of 1–2 % (e.g., Pyle et al., 2018; Cao et al., 2020). This level of intra-laboratory uncertainty is typical for [Ba] data obtained using isotope dilution-inductively coupled plasma mass spectrometry, and applies to GEOTRACES-era datasets and to much of the training data from the Indian Ocean. However, intra-laboratory uncertainty is typically much smaller than interlaboratory uncertainty, which is often between 6–9 % (e.g., Hathorne et al., 2013). At the upper end, the benchmark Ba-Si linear regression achieves a MAPDMAPE of 9.7 % in the Indian Ocean (Fig. 3A4). Thus, useful ML models of [Ba] should achieve uncertainties MAPE between 1–10 %. Indeed, our favored predictor model, #33363080, achieves a MAPDMAPE of 6.30 %.

Now we consider two factors that contribute to the observed 6.30 % uncertainty: realization uncertainty and uncertainties in the training data. The realization uncertainty stems from the fact

that two models trained on the same training dataset—even with the exact same subset of model features—will produce slightly different predictions. This is due to the holdout cross-folding process used during model training, which partitions the training dataset into <u>random</u> subsets (see Sect. 3.1.). Because the partitioning is randomThus, the training process results in a slightly different trained model each time the model is realized. We quantified the realization uncertainty by training model #3336select models 100 times and calculating the relative standard deviation of the different predictions of [Ba] for <u>allthe</u> 3,305,505.3 million values in the output. This uncertainty is small; the median, mean, and maximum realization uncertainty was 0.03 %, 0.04 %, and 0.32 % variability in modeled [Ba].

Next we consider uncertainties in the training data. As noted above, many labs report uncertainties on [Ba] measurements of 1-2 %, while inter-laboratory differences may be up to a factor of five larger. However, this does not consider any uncertainties associated with the other physical and chemical features used to predict [Ba]. In general, these supporting measurement uncertainties should be small-since: all overboard sensors are regularly calibrated and biogeochemical properties in GEOTRACES are determined using established methods that are based on GO-SHIP best practices (Hood et al., 2010). Moreover, all GEOTRACES sections include crossover stations that are intended to facilitate intercalibration of all parameters, including those used here to predict [Ba] (Fig. 2; Cutter, 2013). The WOA, MLD, Chl. a, and bathymetry data products are similarly subjected to stringent quality review and so we consider it unlikely that these data contribute systematic biases. We believe that the most likely source of uncertainty relates to the fact that all predictor information used for model testing in the Indian Ocean was derived from time-averaged data products, whereas [Ba] was derived from *in situ* measurements. We made this decision because the in situ data were incomplete for all 12 core features (Table 1), and this would have necessitated interpolation for some features and not others. Since all models were tested using the same predictor information, the comparison process should avoid systematic errors, though this does not preclude temporal variability, described next.

#### 5.2.3. Other considerations

We now consider four other factors that potentially contribute to the uncertainty of the model output: short- and long-term temporal variations, limitations of ML, and uncertainties regarding the thermodynamic properties of BaSO<sub>4</sub>. Short-timescale variability in [Ba] may affect how models were evaluated, though this effect is difficult to quantify. In principle, the trained models should be able to resolve seasonal variations in [Ba] since they were trained on in situ physical and chemical data. In contrast, model predictions in the Indian Ocean were made using annual average physical and chemical conditions and then evaluated by comparing these predictions against *in* situ [Ba]. The temporal mismatch between Indian Ocean observations and predictions is unlikely to be significant in the deep ocean, where seasonal variations are minor and the Ba residence time is longest (e.g., Hayes et al., 2018). Seasonal variations are, however, likely to matter more for the surface ocean. We were able to minimize some of the impact of these uncertainties by using longterm averages of Chl. a and the maximum monthly mean MLD during model training and testing. Significant seasonal mismatches for other parameters are unavoidable given that [Ba] data are too sparse to develop a time-resolved model. We suspect that these variations are most likely to be significant for boundary sources rather than biogeochemical cycling of Ba; significant biogeochemical drawdown of surface [Ba] over seasonal timescales appears to be rare (e.g., Esser & Volpe, 2002), whereas there are large seasonal variations in river discharge that impact nearshore [Ba] (e.g., Samanta & Dalai, 2016). These suspicions could be tested using a model with better than  $1 \times 1^{\circ}$  spatial resolution, which—in theory—is possible with model #33363080, so long as similarly high-resolution data are provided for the six predictors utilized by this model (z, T, S, z) $[O_2]$ ,  $[PO_4]$ ,  $[NO_3]$ , and  $[NO_3S_i]$ ). While it is challenging to precisely quantify seasonal uncertainties, we note that model #33363080 performs well at low [Ba], which is found mostly near the surface, where seasonal variations should beexhibit the largest (Figs. 3B, 8B).effects. Likewise, seasonal variations will have only a minor effect on our calculations of global mean [Ba] or  $\Omega_{\text{barite}}$  (Fig. 8).

Long-term variability in [Ba] may also influence model performance, since the testing data from the Indian Ocean were collected between 1977 (GEOSECS) and 2008 (SS259; Fig. 2). If secular changes in Indian Ocean [Ba] were occurring, we might expect models to make accurate predictions for some datasets at the expense of others. In contrast, we note that model #33363080

reproduces all testing datasets similarly well, with the exception of a subset of samples from SS259 in the deep Bay of Bengal (Fig. 3C). Here we observe that model #3336 tends to predict -93080 predicts 18 % higher [Ba] than observed by Singh et al. (2013), particularly around 2) for the 42 samples between 1,000-3,000 m (Fig. 6AFigs. 4B; 7A, B). HoweverInterestingly, model #33363080 correctly predicts [Ba] at nearby GEOSECS stations 445 and 446, also in the Bay of Bengal, sampled some 31 years prior. The to SS259. We briefly consider three possibilities for the origin of the this regional model-data discrepancy-. It may derive from the fact that model #3080 does not include the features needed to correctly predict [Ba] in this region these samples. We view this as the least likely possibility as model #3080 performs well for other samples from the northern Indian Ocean, including samples shallower than 1,000 m from Singh et al. (2013). Another possibility is uncertain; we speculate that it may reflect could reflect an 18 % decrease in [Ba] in the deep Bay of Bengal since the GEOSECS survey in the 1970's. Lastly, it could reflect differences in how in situ [Ba] was measured, noting that Singh et al. (2013) opted for standard addition overinstead of isotope dilution. Alternatively, it could reflect a ~9 % decrease in [Ba] in the deep Bay of Bengal since the 1970's. We currently lack the data needed to confidently distinguish between these latter two possibilities.

A third factor concerns the limitations of ML itself. We note that no trained model was able to achieve a MAPDMAPE better than ~6 %. This <u>6 %</u> value may represent one of three things. First, it may point toward an intrinsic limitation of Gaussian Process Regression. Other types of ML, such as Decision Trees or Artificial Neural Networks, may be able to achieve superior accuracy, though this was not investigated. Second, it may indicate that the 12 features investigated provide insufficient information about [Ba] to achieve higher accuracy. We view this as unlikely given that our earlier analysis showed that only <u>5 -8six-nine</u> features were needed to accurately simulate [Ba] and that the 12 features investigatedtested have proved useful in other studies simulating dissolved tracer distributions (e.g., Rafter et al., 2019; Sherwen et al., 2019; Roshan & DeVries, 2021). However, this does not rule out the existence of other features beyond the 12 that we tested that are more useful for predicting [Ba], only that we did not investigate them. Third, it is possible that the lowest MAPDMAPE of ~6 % reflects the current limit of inter-laboratory uncertainty in determining [Ba]. We note that inter-laboratory uncertainties of 6–9 % were reported for the measurement of Ba:Ca in carbonates (n = 10 labs; Hathorne et al., 2013). If the ~6 % MAPDMAPE

derives from inter-laboratory uncertainty, it is unlikely that further model refinements will improve the accuracy of [Ba] predictions: the fundamental limitation is the data, not the model.

A final source of uncertainty concerns the computation of  $\Omega_{\text{barite}}$ , which contains two further sources of uncertainty: the thermodynamic model and the solubility coefficients used to calculate  $\Omega_{\text{barite}} \underline{K}_{\text{sp.}}$ . We calculated  $\Omega_{\text{barite}}$  based on the computation described by Rushdi et al. (2000), and our approach yields similar values to their study and several others (e.g., Jeandel et al., 1996;(2000). This calculation Monnin et al., 1999; see Appendix). The model used by Rushdi et al. (2000) is based on BaSO<sub>4</sub> solubility data from Raju & Atkinson (1988), who note good agreement with the thermodynamic data of Blount (1977). These solubility data were obtained based on experimentation with lab-made, coarse-grained BaSO<sub>4</sub>, which is unlikely to be wholly representative of the microcrystalline BaSO<sub>4</sub> precipitates found in seawater. Thus, the absolute values of  $\Omega_{\text{barite}}$  calculated here may be subject to eventual revision; however, the vertical (Fig. 1), spatial (Figs. 3-74-8), and whole-ocean (Fig. 89) trends in  $\Omega_{\text{barite}}$  are robust. Should new thermodynamic data for marine-relevant micron-sized pelagic BaSO<sub>4</sub> become available, updated maps of  $\Omega_{\text{barite}}$  could be easily recalculated using existing model #3336-derived [Ba] data. recalculated using model #3080-derived [Ba] data. Given the nature of these uncertainties, we opted to calculate prediction uncertainties for  $\Omega_{\text{barite}}$  empirically by comparison to literature data (see Appendix). This yields a value between 0.08 and 0.10, similar to the 10 % prediction error reported by Monnin et al. (1999).

We can calculate  $\Omega_{\text{barite}}$  to a high degree of precision; however, there are numerous uncertainties pertaining to ML-predicted [Ba], the BaSO<sub>4</sub> solubility coefficients used to calculate  $K_{\text{sp}}$ , and the thermodynamic model used in the computation of  $\Omega_{\text{barite}}$  (Sect. 5.2.). Thus,

# 5.3. Barium in seawater: A global perspective Model applications

Here we provide an overview of the main model features in [Ba], Ba\* and  $\Omega_{\text{barite}}$ , then outline fourthree possible applications of the model output. Predictions from model #3336 shows

#### 5.3.1. Dissolved distribution of [Ba]

<u>Model #3080 predictions show</u> several interesting features in [Ba] (Figs. 4-7). <u>Model #33365-8)</u>. <u>The model</u> reproduces the <u>expected</u> nutrient-like distribution of [Ba] (Fig. 1C) and shows a general increase in [Ba] along the Meridional Overturning Circulation; <u>volume-weighted</u> mean [Ba] increases from 67 to <u>8890</u> to 106 nmol kg<sup>-1</sup> from the Atlantic to Indian to the Pacific Ocean, respectively. The model also predicts some variation in shallow [Ba] that follows major surface-water currents, such as a region of elevated [Ba] associated with the North Pacific Current, as well as low [Ba] in the western North Atlantic associated with the Gulf Stream (Fig. <u>4B5B</u>; Talley et al., 2011). <u>TakingHowever</u>, these features and the processes driving them await corroboration.

<u>Considering</u> the ocean as a whole, we can <u>also</u>-use our model <u>output</u> to calculate the total Ba inventory of <u>the oceanseawater</u>. Using the mean oceanic [Ba] of 89 nmol kg<sup>-1</sup> and multiplying by the mass of seawater  $(1.37 \times 10^{21} \text{ kg})$  yields a total inventory of  $122\pm 87$  Tmol Ba, whereby the <u>rangeuncertainty</u> is based on the <u>MAPDMAPE</u> of model <u>#33363080</u> (6.30 %). This estimate of the total oceanic Ba inventory is <u>approximately 10between 11</u>–21 % lower than <u>previousexisting</u> estimates of 145 Tmol Ba (Dickens et al., 2003; Carter et al., 2020). <u>Thus, given current estimates</u> <u>Given the range of probable global marine Ba fluxes of between 18 (Paytan & Kastner, 1996) and</u> 44 Gmol Ba yr<sup>-1</sup> (Rahman et al., 2022), <u>our inventory estimate places</u> the mean residence time of Ba in seawater <u>is likely</u> between 2,700600–7,200 years.

#### Next we consider the main features in

# 5.3.2. The Ba\*\_Si relationship

We now quantify spatial and  $\Omega_{\text{barite}}$ -vertical variations in the Ba–Si relationship, which we explore using Ba\*. Star tracers, such as Ba\*, highlight the processes affecting the distribution of a tracer by comparing it to another tracer that shares the same circulation (Gruber & Sarmiento, 1997). The concept has since been extended to study the processes affecting the distributions of many other bioactive elements, including Si (Si\*, relative to N; Sarimento et al., 2004), cadmium (Cd\*, relative to P; Baars et al., 2014), zinc (Zn\*, relative to Si; Wyatt et al., 2014). First defined by Horner et al. (2015) for Ba, Ba\* is analogous to other star tracers: it is a measure of Ba–Si decoupling whereby larger values indicate larger Ba–Si deviations relative to expected mean ocean behavior. Vertical or spatial differences in Ba and Si sources or sinks will drive variations in Ba\*, as will any Ba:Si fractionation occuring during their combined cycling. Conversely, if all Ba and Si cycling occurs in the same places (and with a fixed Ba:Si ratio), no Ba–Si decoupling will occur and Ba\* will exhibit conservative behavior. Since Ba and Si are cycled by different processes *and* there are large vertical and spatial variations in the intensity of these processes (e.g., Bishop, 1989), significant variations in Ba\* are possible. We now explore these variations.

In the surface ocean, patterns of Ba\* generally resemble those of [Ba] (Fig. 4). This is likely because inIn large parts of the ocean, surface [Si] approaches 0 µmol kg<sup>-1</sup>; thus, variations in Ba\* derive mostly from variations in [Ba]. Barium-star is, however, strongly positive in the top 200 m of the This is most evident when examining regions with significant terrestrial input of Ba, such as from major rivers (Sect. 5.2.1) and from rivers and continental shelves in the Arctic (e.g., Guay & Falkner, 1998; Whitmore et al., 2022; Fig. 5A). The Southern Ocean, even also exhibits positive Ba\*, though [Si] is in the 10's of µmol kg<sup>-1</sup> we suspect the mechanism is different. Here we observe a belt of waters with positive Ba\* ≈+20 nmol kg<sup>-1</sup> range. centered on the Polar Frontal Zone—the region between the Antarctic Polar Front and the Subantarctic Front (Orsi et al., 1995; Fig. 5A). Silicic acid is intensely stripped from waters that transit northward through this region (e.g., Sarmiento et al., 2004), potentially contributing to elevated Ba\* at the sea surface. Dissolved [Ba] and Ba\* then decrease to the north of the Subantarctic front, partly driven by extensive particulate Ba formation in the frontal region (e.g., Bishop, 1989).

At 1,000 m, the Atlantic, South Pacific, and southern Indian Oceans exhibit positive Ba\* around  $\pm 10 \text{ nmol } \text{kg}^{-1}$ , whereas the North Pacific, Southern, and Indian Oceans exhibit negative values between  $\pm 10 \text{ to} \pm 20 \text{ nmol } \text{kg}^{-1}$ , whereas the Atlantic and South Pacific northern Indian Oceans are positive around  $\pm 10 \text{ nmol } \text{kg}^{-1}$  (Fig. 5). Below, 1,000 m, the Southern and negative between  $\pm 10 \text{ nmol } \text{kg}^{-1}$  (Fig. 5). Below, 1,000 m, the Southern and negative between  $\pm 10 \text{ nmol } \text{kg}^{-1}$  (Fig. 6C). The positive anomalies are likely related to the northward spreading of southern-sourced intermediate waters that originate within the Polar Frontal Zone and carry positive Ba\* into the low latitudes (e.g., Bates et al., 2017). In the Atlantic Oceans do not exhibit significant gradients in Ba\* and are, these values are carried all the way to the north of the basin and return as North Atlantic Deep Water with only minor modifications to Ba\* ( $\approx \pm 10 \text{ nmol } \text{kg}^{-1}$ ; Figs. 6C, 7C, 8C). Negative Ba\* in the North Pacific, Southern, and northern Indian Ocean at 1,000 m likely reflects a mixture of hydrographic processes and *in situ* processes. For example, the extensive region of negative and positive downBa\* in the North Pacific is closely associated

with North Pacific Intermediate Water, which originates in the Sea of Okhotsk (Talley, 1991). While the specific mechanism sustaining this particular Ba\* feature is unknown, it most possibly reflects a combination of preferential removal of Ba relative to Si in the source water formation region (such as from particulate Ba formation) and weak vertical mixing in the subsurface North Pacific relative to lateral transports (e.g., Kawabe & Fujio, 2010). We suspect that the negative Ba\* values seen above 1,000 m in the northern Indian Ocean originate through processes occurring internally within this basin, as the majority of the Indian Ocean below 1,000 m exhibits positive Ba\*. A possible mechanism for these shallow negative Ba\* anomalies may relate to the relatively weak overturning transports (Talley, 2008) and strong particulate Ba cycle north of 30 °S (Singh et al., 2013), though this awaits more detailed investigation.

Lastly, the Southern Ocean exhibits negative Ba\* between -10 and -20 nmol kg<sup>-1</sup> from  $\approx 200$  m water depth to the seafloor, respectively. In contrast, the Indian and North Pacific Oceans. These negative anomalies in Ba\* appear to be associated with Circumpolar Deep Water and, below that, Antarctic Bottom Water; the influence of the latter can also be seen in near-bottom negative Ba\* in the South Pacific, southern Indian, and South Atlantic Oceans (Fig. 8C). As with the other basins, the origin of the negative Ba\* waters in the Southern Ocean likely reflects a combination of *in situ* and circulation-related phenomena. For example, in the Southern Ocean, Si is only stripped at the very surface, whereas particulate Ba formation is thought to be greatest in the mesopelagic (i.e., between 200-1,000 m; e.g., Stroobants et al. 1991). Barite formation is generally considered to be related to the regeneration of particulate organic matter (e.g., Chow & Goldberg, 1960), whereby the former consumes Ba and the latter releases Si. Thus, intense organic matter remineralization and associated pelagic BaSO<sub>4</sub> precipitation could contribute to negative Ba\* in the mesopelagic Southern Ocean. Similarly, the Si cycle in the Southern Ocean tends to 'trap' a significant fraction of the global Si inventory in the waters circulating close to Antartica (e.g., Holzer et al., 2014). Since the calculation of Ba\* depends on both [Ba] and [Si], waters with elevated [Si] will exhibit positive Ba\* around 2,000 m, between +5 and +15 nmol kg<sup>-1</sup> (Fig. 6). The most positive Ba\* values, between +20 and +30 nmol kg<sup>-1</sup>, are observed at lower Ba\* whether or not there is elevated Ba removal.

By 2,000 m, almost all of the ocean north of 50 °S exhibits positive Ba\* (Fig. 7C). By 4,000 m in the Pacific, specifically in the, the areal extent of the positive-Ba\* waters shrinks to encompass

the area north of 30 °S (Fig. 8C). Despite covering a smaller area, the abyssal ocean exhibits the most positive Ba\* values outside of the surface of the Southern Ocean. The reasons for elevated and increasing Ba\* between the deep and abyssal oceans likely reflects a mixture of local and regional processes, and we offer two speculative explanations for these patterns. First, Si trapping in the Southern Ocean potentially renders most of the deep ocean away from Antarctica deficient in Si relative to Ba. Thus, much of the ocean may exhibit more positive Ba\* than the deep circum-Antarctic region due to processes unrelated to Ba cycling. Second, the most positive Ba\* values are generally found close to the seafloor, rather than the mid-depths, especially in the North Pacific, the Peru and Chile Basins as well as, and the Philippine Sea (Fig. 7). Vertical profiles of  $\Omega_{barite}$  are. This may indicate a mechanism that preferentially removes Ba (relative to Si) from the mid-depths, or input of Ba (relative to Si) close to the seafloor.

Systematic variations in Ba\* arise due to differences in the marine biogeochemical cycles of Ba and Si. While, in some cases, the specific drivers of these variations remains unresolved, our model identifies multiple hotspots of Ba–Si decoupling that warrant additional study.

# 5.3.3. Barite saturation state of seawater

Here we show that our approach can predict  $\Omega_{\text{barite}}$  with an MAE of 0.08, that our output is in agreement with published values; comparisons are provided in the Appendix, and that the deep ocean, below 1,000 m, is at saturation with respect to BaSO<sub>4</sub>. By comparison to literature data, we estimate that our model achieves a typical prediction uncertainty on  $\Omega_{\text{barite}}$  of 0.08 (see Appendix). Accordingly, values of  $\Omega_{\text{barite}}$  between 0.92–1.08 can be considered as 'BaSO<sub>4</sub> saturated,' whereas values of  $\Omega_{\text{barite}}$  <0.92 or >1.08 indicate under- or super-saturation, respectively. Global patterns in  $\Omega_{\text{barite}}$  derived using our model are similar to those reported by (e.g., Jeandel et al., 1996; Monnin et al., \_\_(1999;) and Rushdi et al., \_\_(2000). Excepting the Readers looking for detailed basin-bybasin descriptions of  $\Omega_{\text{barite}}$  are directed to those studies. Briefly our model shows that, excepting the high latitudes, the surface ocean is undersaturated with respect to BaSO<sub>4</sub> (i.e.,  $\Omega_{\text{barite}} <1$ ) and the(0.92). The lowest values of  $\Omega_{\text{barite}}$  in the open ocean are observed in the hot, salty cores of the Subtropical Gyres ( $\Omega_{\text{barite}}$  between 0.1–0.2; Figs. 4D, 8D). WhereasFig. 5D). Conversely, the cold and fresh polar regions exhibit supersaturation at the sea surface, though there are important differences between the Southern and Arctic Oceans. The Southern Ocean remains supersaturated (i.e.,  $\Omega_{barite} > 1$ )exhibits BaSO<sub>4</sub> saturation to depths around 2,000 m, whereas the Arctic Ocean switches to undersaturated conditions below ~the halocline (~250 m. Below). At 1,000 m, most of the North Pacific is supersaturated achieves saturation (or slight supersaturation) with respect to BaSO<sub>4</sub> (Fig. 6D) and byat 2,000 m almost all of the ocean exhibits  $\Omega_{barite} > 1$ , excepting 0.92. The main exceptions to this are the Atlantic Ocean, which is undersaturated at all depths, and the southern Indian Ocean between 35–50 °S (Fig. 6D7D). The South Pacific and Indian Oceans return to undersaturated conditions by 4,000 m, whereas the majorityparts of the North Pacific exhibits  $\Omega_{barite} > 1$  remain saturated to the seafloor (Fig. 7D8D). From a volumetrieglobal perspective, the oceans are slightly undersaturated with respect to BaSO<sub>4</sub>: volume-weighted mean  $\Omega_{barite} = 0.82$ ; however, the ocean belowbetween 1,000–5,500 m exhibits a mean  $\Omega_{barite} \ge 0.92$ , which (Fig. 9). This result implies that much of the deep ocean, as a whole, is close to saturationchemical equilibrium with respect to BaSO<sub>4</sub> (Fig. 8D).

#### 5.3.4. Model applications Lastly, in

In the spirit of maximizing model utility, we suggest three possible uses for these datamodel #3080 outputs. First, the outputs can be used for model intercomparison and intercalibration. For example, a number of statistical models, such as Optimum Multiparameter Optimization, have been successfully used to study Ba cycling in the North Atlantic (Le Roy et al., 2018; Rahman et al., 2022), Southeast Pacific (Rahman et al., 2022), and Mediterranean Sea (Jullion et al., 2017). These models can apportion the relative contributions of *in situ* biogeochemical cycling and conservative mixing to observed [Ba]; however, accurate quantification of these processes requires *a priori* knowledge of end-member water mass [Ba], which model #33363080 can provide. Our model could also be used to benchmark output from process-based models, such as the-Ocean Circulation Inverse ModelModels (e.g., John et al., 2020; Roshan & DeVries, 2021). Second, the output can be used for interpolation purposes. For example, manyMany groups investigated Ba partitioning into various types of marine carbonates (see Sect. 1 for examples); however, these investigations are sometimes performed without a co-located measurement of [Ba]. In these cases output from model #33363080 could be used to help calibrate specific substrates, such as deep-sea corals or benthic forams. This also avoids the potential for circular reasoning whereby [Si] is used to estimate [Ba], which is then reconstructed from the Ba:Ca ratio of carbonates to estimate [Si]. Third, the model output makes testable predictions for regions of the ocean that have yet to be sampled by GEOTRACES-style surveys. Several of these regions, such as the Southern Ocean, exhibit with sharp lateral and vertical gradients in [Ba], Ba\*, and  $\Omega_{\text{barite}}$ . Such gradients should be considered prime targets for future process-oriented studies of marine Ba cycling.

# 6. Data availability

Data described in this manuscript can be accessed at the *Biological and Chemical Oceanography Data Management Office* under data doi:10.26008/1912/bco-dmo.885506.42 (Horner & Mete, 2023).

# 7. Conclusions

This study presents a spatially and vertically resolved global model of [Ba] determined using Gaussian Process Regression machine learning. The model reproduces several known features of the marine [Ba] distribution and makes testable predictions in regions that are yet to be sampled. Analysis of the model output reveals the mean oceanic [Ba] is 89 nmol kg<sup>-1</sup>, implying a total marine Ba inventory of  $122\pm87$  Tmol. Using predictors from the World Ocean Atlas, we also estimate the global distribution of Ba\* and  $\Omega_{\text{barite}}$ . Both properties exhibit significant gradients that can be systematically investigated in future studies. The mean oceanic  $\Omega_{\text{barite}}$  is 0.82, though between 1,000–5,500 m the mean is  $\geq 0.92$ , implying that the deep ocean is elose to saturationat equilibrium with respect to BaSO4. Our model output should prove valuable in studies of Ba biogeochemistry, specifically for statistical- and process-based model validation, calibrating sedimentary archives, and for identifying promising regions for further study. More broadly, our study demonstrates the utility of using machine learning to accurately simulate the distributions of trace elements in seawater. With minor adjustments, our approach could be employed to make predictions for other dissolved tracers in the sea.

# Appendix

Here we compare our results with published profiles of  $\Omega_{\text{barite}}$ . Our results were calculated using the thermodynamic model of Rusdi et al. (2000), model #33363080 [Ba], and WOA *T*, *S*, and pressure. Literature profiles of  $\Omega_{\text{barite}}$  were calculated using one of three different thermodynamic models and *in situ* observations of [Ba], *T*, *S*, and pressure. In general, there is strong agreement between modeled and *in situ*  $\Omega_{\text{barite}}$  whereby our model reproduces the shape of published profiles (Fig. A1). There are, however, some small systematic offsets between the various approaches, and we suspect that these derive from differences in the underlying thermodynamic models.



Figure A1. Comparison of literature- (symbols) and Model #33363080-derived (dashed line) estimates values of  $\Omega_{\text{barite}}$ . Panels A and B show profiles of  $\Omega_{\text{barite}}$  at GEOSECS St. 89 (60°0' S, 0°2' E). The other panels are from the Indian Ocean; C and D are from INDIGO 2 St. 36 (6°9' S, 50°55' E) and E from GEOSECS St. 420 (0°3' S, 50°55' E), some ≈675 km north of INDIGO 2 St. 36.

We compare our model output with literature data  $\Omega_{\text{barite}}$  at two stationslocations in two basins (Fig. A1). These stationslocations were selected for comparing  $\Omega_{\text{barite}}$  because chosen to ensure a fair comparison between studies; at both locations each location, at least two studies calculated profiles of  $\Omega_{\text{barite}}$  using the same underlying *in situ* data for [Ba], *T*, *S*, and pressure. This ensures a fair comparison between studies, since Thus, any differences in modeled  $\Omega_{\text{barite}}$  should derive from the thermodynamic model and not the input data. Likewise, the literature profiles at these locations were based on calculations for pure, rather than strontian, BaSO<sub>4</sub>, as in our study. Published profiles of  $\Omega_{\text{barite}}$  were extracted graphically from each study using *WebPlotDigitizer*  (Rohatgi, 2022). This extraction process may introduce some minor scatter in the literature data, though this is relatively minor relative<u>compared</u> to the range of variation in  $\Omega_{\text{barite}}$ .

First, we examine profiles of  $\Omega_{\text{barite}}$  reported for GEOSECS St. 89 in the Southern Ocean (Fig. A1; Monnin et al., 1999; Rushdi et al., 2000). Modeled and published profiles show supersaturation in the surface ocean and undersaturation below 2,000–2,500 m. Profiles from Rushdi et al. (2000) show excellent agreement with  $\Omega_{\text{barite}}$  calculated from model #33363080 [Ba] and WOA *T*, *S*, and pressure, with our output slightly negatively offset by a MADMAE of 0.06 (n = 22). Given that we use the same thermodynamic model as Rushdi et al. (2000), the overall excellent agreement with their study is not surprising. However, the result is nonetheless reassuring since our study uses mean annual values for the various inputs, whereas Rushdi et al. (2000) utilized *in situ* data. There is a slightly larger offset between our profile of  $\Omega_{\text{barite}}$  and that calculated by Monnin et al. (1999), with our respective profile offset to higher  $\Omega_{\text{barite}}$  by a MADexhibiting an MAE of 0.13 (n = 41). This most likely reflects differences in the underlying thermodynamic model and not the *in situ* data since our model reproduces the same overall profile shape as Monnin et al. (1999). Likewise, both Monnin et al. (1999) and Rushdi et al. (2000) used the same *in situ* input data and their results are highly comparable, albeit with an offset similar to that between our results and Monnin et al. (1999).

Next we examine profiles of  $\Omega_{\text{barite}}$  in the Indian Ocean for samples from INDIGO 2 St. 36 (Fig. A1; Jeandel et al., 1996; Rushdi et al., 2000). Profiles of  $\Omega_{\text{barite}}$  show undersaturation at the surface, moderate supersaturation between 2,000–3,500 m, then return to undersaturated conditions down to the seafloor. Our profile shows overall excellent agreement with that of Jeandel et al. (1996), whereby our data are offset to slightly lower a comparison of  $\Omega_{\text{barite}}$  withyields a MADMAE of of 0.0603 (n = 21). The largest differences are observed between 594–1,042 m depth, where the MAD is 0.15 (n = 3). Our profile shows similarly good agreement with that of Rushdi et al. (2000), whereby our data are offset to lower a comparison between our respective values of  $\Omega_{\text{barite}}$  withyields a MADMAE of 0.0704 (n = 20). As with the data of Jeandel et al. (1996), we observe a larger offset between modeled  $\Omega_{\text{barite}}$  and the data of Rushdi et al. (2000) between 594–1,042 m, equivalent to a MAD of 0.17 (n = 3). We consider these larger mesopelagic offsets in  $\Omega_{\text{barite}}$  to be the result of differences in the predictors (i.e., [Ba], *T*, *S*), rather than the thermodynamic model;

Jeandel et al. (1996) and Rushdi et al. (2000) use the same *in situ* predictor data and yield similar  $\Omega_{\text{barite}}$ , despite using different thermodynamic models.

We also compared our results with data from St. 420 of GEOSECS (Monnin et al., 1999), which is located  $\approx 675$  km north of INDIGO 2 St. 36 (Fig. 2). As with data from the Southern Ocean (GEOSECS St. 89), our profile data are offset to higher  $\Omega_{\text{barite}}$  than those of Monnin et al. (1999) by a MAD), with slightly larger MAE of 0.1216 (n = 29). However, our modeled  $\Omega_{\text{barite}}$  is generally in much closer agreement with Monnin et al. (1999) above 1,250100 m than below, equivalent to a MADMAE of 0.0304 (n = 98) and 0.1621 (n = 2021), respectively. In this case it is more challenging to ascribe a unique cause of the differences in calculated  $\Omega_{\text{barite}}$ ; these offsets could relate to differences in the predictors or the thermodynamic model.

We can use these comparisons to estimate the prediction uncertainty on our model-derived values of  $\Omega_{\text{barite.}}$  The MAE of the 133 comparisons shown in Fig. A1 yields a value of 0.10. However, there are different numbers of points in each profile; we thus believe it is more appropriate to average the MAE calculated for each of the five profiles, which yields a value of 0.08. Both values are similar to the 10 % prediction uncertainty reported by Monnin et al. (1999).

Overall, our ML-derived profiles of  $\Omega_{\text{barite}}$  show excellent agreement with *in situ* data, both in terms of profile shape and absolute-values of  $\Omega_{\text{barite}}$ . We use this comparison to within estimate the prediction uncertainty on ML-derived values of  $\Omega_{\text{barite}}$ , which we calculate as being between 0.108 and 0.10. Should a revised thermodynamic model and/or improved BaSO<sub>4</sub> solubility coefficients become available, a new grid of  $\Omega_{\text{barite}}$  could be calculated using Model #33363080 [Ba] and WOA *T*, *S*, and pressure data.

# Author contributions

Project conceptualization and funding acquisition by T.J.H. Data curation, formal analysis, investigation, and methodology by O.Z.M., A.V.S., <u>H.H.K.</u> and T.J.H. Data visualization by A.V.S. and T.J.H. Software provided by O.Z.M., A.V.S., H.H.K., and A.G.D. Writing (original draft) by O.Z.M. and T.J.H.; review and editing by <u>A.V.S.</u>, H.H.K., A.G.D., L.M.W., A.M.S., M.G., and W.D.L.

# **Competing interests**

The authors declare that they have no conflict of interest.

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