**Deep-sea sediments of the global ocean**

Markus Diesing

1Geological Survey of Norway (NGU), Postal Box 6315 Torgarden, 7491 Trondheim, Norway

Correspondence to: Markus Diesing (markus.diesing@ngu.no)

**Abstract.** Although the deep-sea floor accounts for approximately 60% of the Earth’s surface, there has been little progress in relation to deriving maps of seafloor sediment distribution based on transparent, repeatable, and automated methods such as machine learning. A new digital map of the spatial distribution of seafloor lithologies below 500 m water depth is presented to address this shortcoming. The lithology map is accompanied by estimates of the probability of the most probable class, which may be interpreted as a spatially-explicit measure of confidence in the predictions, and probabilities for the occurrence of five lithology classes (Calcareous sediment, Clay, Diatom ooze, Lithogenous sediment, and Radiolarian ooze). These map products were derived by the application of the Random Forest machine learning algorithm to a homogenised dataset of seafloor lithology samples and global environmental predictor variables that were selected based on the current understanding of the controls on the spatial distribution of deep-sea sediments. It is expected that the map products are useful for various purposes including, but not limited to, teaching, management, spatial planning, design of marine protected areas and as input for global spatial predictions of marine species distributions and seafloor sediment properties. The map products are available at https://doi.org/10.1594/PANGAEA.911692 (Diesing, 2020).

**1 Introduction**

The deep-sea floor accounts for >85% of seafloor area (Harris et al., 2014) and >60% of the Earth’s surface. It acts as a receptor of the particle flux from the surface layers of the global ocean, is a place of biogeochemical cycling (Snelgrove et al., 2018), records environmental and climate conditions through time and provides habitat for benthic organisms (Danovaro et al., 2014). Being able to map the spatial patterns of deep-sea sediments is therefore a major prerequisite for many studies addressing aspects of marine biogeochemistry, deep-sea ecology, and palaeo-environmental reconstructions.

Until recently, maps of global deep-sea sediments were essentially variants of a hand-drawn map presented by Berger (1974) and typically depicted five to six sediment types, namely calcareous ooze, siliceous ooze (sometimes split into diatom ooze and radiolarian ooze), deep-sea (abyssal) clay, terrigenous sediment and glacial sediment. Since then, Dutkiewicz et al. (2015) collated and homogenised approximately 14,500 samples from original cruise reports and interpolated them using a support vector machine algorithm (Cortes and Vapnik, 1995). Their map displayed the spatial distribution of 13 lithologies across the world ocean and exhibited some marked differences from earlier maps.
The controls on the distribution of deep-sea sediments have long been discussed (e.g., Seibold and Berger, 1996): Biogenous oozes (>30% microscopic skeletal material by weight) dominate on the deep-sea floor and their composition is controlled by productivity in overlying surface ocean waters, dissolution during sinking and sedimentation and dilution with other materials. The ocean is undersaturated with silica. Preservation of siliceous shells is therefore a function of shell thickness, sinking time (water depth) and water temperature, as siliceous shells dissolve slower in colder water. The dissolution of calcareous shells is increased with increasing pressure (water depth) and CO₂ content of the water (decreasing temperature). The water depth at which the rate of supply with calcium carbonate to the sea floor equals the rate of dissolution (calcite compensation depth; CCD) varies across ocean basins. Deep-sea clays dominate in the deepest parts of ocean basins below the CCD. Deposition of terrigenous material is thought to be a function of proximity to land (distance to shore).

Dutkiewicz et al. (2016) investigated the bathymetric and oceanographic controls on the distribution of deep-sea sediments with a quantitative machine-learning approach. The influence of temperature, salinity, dissolved oxygen, productivity, nitrate, phosphate, silicate at the sea surface and bathymetry on lithogenous sediment, clay, calcareous sediment, radiolarian ooze and diatom ooze were quantified. They found that bathymetry, sea surface temperature and sea surface salinity had the largest control on the distribution of deep-sea sediments. Calcareous and siliceous oozes were not linked to high surface productivity according to their analysis. Diatom and radiolarian oozes were associated with low sea surface salinities and discrete sea surface temperature ranges.

The aim of this study is to derive a map of deep-sea sediments of the global ocean by utilising environmental predictor variables for the development and application of a machine-learning spatial prediction model. Besides a categorical map giving the spatial representation of seafloor types in the deep sea, probability surfaces for individual sediment classes and a map displaying the probability of the most probable class in the final prediction will also be provided.

2 Data

2.1 Predictor variables

The initial choice of the predictor variables was informed by the current understanding of the controls on the distribution of deep-sea sediments and the availability of data with full coverage of the deep sea at a reasonable resolution. We chose predictor variables mentioned above, but also included sea-surface iron concentration, which was not available to Dutkiewicz et al. (2016), but is an important nutrient for phytoplankton (Table 1). The predictor variable raster layers from Bio-ORACLE (Assis et al., 2018; Tyberghein et al., 2012) and MARSPEC (Sbrocco and Barber, 2013) were utilised. Whenever available, statistics of the variable other than mean were downloaded. These included the minimum, maximum and the range (maximum – minimum).
2.2 Response variable

The response variable is seafloor lithology, a qualitative multinomial variable. The seafloor sediment sample data (seafloor_data.npz) from Dutkiewicz et al. (2015) were downloaded from ftp://ftp.earthbyte.org/papers/%20Dutkiewicz_etal_seafloor_lithology/iPython_notebook_and_input_data/. The original dataset consisted of 13 seafloor lithology classes, while Dutkiewicz et al. (2016) simplified these to five major classes. The latter scheme was chosen here (Table 2), as the five major classes agree well with lithologies typically depicted in hand-drawn maps. For a detailed description of the original lithology classes refer to GSA Data Repository 2015271 (https://www.geosociety.org/datarepository/2015/2015271.pdf).

3 Methods

The general workflow for building a predictive spatial model was outlined by Guisan and Zimmermann (2000). This involves five main steps: (1) Development of a conceptual model, (2) statistical formulation of the predictive model, (3) calibration (training) of the model, (4) model predictions and (5) evaluation of the model results (accuracy assessment). The conceptual model was already presented in the introduction. The remaining steps are described in the following sections. The analysis was performed in R 3.6.1 (R Core Team, 2018) and RStudio 1.2.1335 and is documented as an Executable Research Compendium (ERC), which can be accessed at https://o2r.uni-muenster.de/#/erc/GWME2voTDdb5oeaQFuTWMCEMveKS1MiXm.

3.1 Data pre-processing

The raster layers (predictor variables) were stacked, limited to water depths below 500 m, projected to Wagner IV global equal-area projection with a pixel resolution of 10 km by 10 km and scaled. The sample data (response variable) were pre-processed in the following way: Only samples of the five major lithologies (Table 2) deeper than 500 m were used and duplicates were removed from the original sample dataset. The number of records was therefore reduced from 14,400 to 10,438. The data were projected to Wagner IV. Locations of the sample locations and their respective lithology class are shown in Fig. 1. Predictor variable values were extracted for every sample location. The class frequencies are shown in Table 2.

3.2 Predictor variable selection

Variable selection reduces the number of predictor variables to a subset that is relevant to the problem. The aims of variable selection are three-fold: (1) to improve the prediction performance, (2) to enable faster predictions and (3) to increase the interpretability of the model (Guyon and Elisseeff, 2003). It is generally advisable to reduce high-dimension datasets to uncorrelated important variables (Millard and Richardson, 2015). Here, a two-step approach was utilised to achieve this goal.
The first step identifies those variables that are relevant to the problem. The second step minimises redundancy in the remaining predictor variables.

Initially, the Boruta variable selection wrapper algorithm (Kursa and Rudnicki, 2010) was employed to identify all potentially important predictor variables. Wrapper algorithms identify relevant features by performing multiple runs of predictive models, testing the performance of different subsets (Guyon and Elisseeff, 2003). The Boruta algorithm creates so-called shadow variables by copying and randomising predictor variables. Variable importance scores for predictor and shadow variables are subsequently computed with the Random Forest algorithm (see below). The maximum importance score among the shadow variables (MZSA) is determined and for every predictor variable, a two-sided test of equality is performed with the MZSA. Predictor variables that have a variable importance score significantly higher than the MZSA are deemed important, while those with a variable importance score significantly lower than the MZSA are deemed unimportant. Tentative variables have a variable importance score that is not significantly different from the MZSA. Increasing the maximum number of iterations (maxRuns) might resolve tentative variables (Kursa and Rudnicki, 2010). Only important variables were retained for further analysis.

The Boruta algorithm is an “all-relevant” feature selection method (Nilsson et al., 2007), which identifies all predictors that might be relevant for classification (Kursa and Rudnicki, 2010). It does not address the question of redundancy in the predictor variable data, which would be required for “minimal optimal” feature selection (Nilsson et al., 2007) usually preferred for model building. To limit redundancy, a second step seeks to identify predictor variables that are correlated with other predictors of higher importance. To achieve this, the Boruta importance score was used to rank the remaining predictor variables. Beginning with the most important variable, correlated variables with lower importance were subsequently removed. Values of the correlation coefficient $r$ were trialled between 0.1 and 1 with a step size of 0.01 to find an appropriate $r$ value that strikes a balance between prediction performance and model interpretability.

### 3.3 Environmental space

It is generally preferable to apply a suitable sampling design for model calibration and evaluation. This would ensure that the environmental variable space is sampled in a representative way. Various methods have been proposed to optimise sampling effort, including stratified random, generalised random tessellation stratified (Stevens Jr and Olsen, 2003) and conditioned Latin hypercube sampling (Minasny and McBratney, 2006) among others. However, such approaches are not feasible here due to time and financial constraints. Instead, we utilised available (legacy) sampling data. It might nevertheless be prudent to assess to what extent the selected samples cover the environmental space of the predictor variables. This was achieved by creating a random subsample ($n = 10,000$) of the selected environmental predictor variables and displaying the density distribution of the random subsample together with the density distribution of environmental variables based on the observations. This allows for a qualitative check to what degree the environmental space is sampled in a representative way.
3.4 Random Forest classification model

The Random Forest (RF) prediction algorithm (Breiman, 2001) was chosen for the analysis due to its high predictive performance in a number of domains (Che Hasan et al., 2012; Cutler et al., 2007; Diesing et al., 2017; Diesing and Thorsnes, 2018; Huang et al., 2014; Prasad et al., 2006). The RF is an ensemble technique based on classification trees (Breiman, 1984). Randomness is introduced in two ways: by constructing each tree from a bootstrapped sample of the training data, and by using a random subset of the predictor variables at each split in the tree growing process. As a result, every tree in the forest is unique. By aggregating the predictions over a large number of uncorrelated trees, prediction variance is reduced and accuracy improved (James et al., 2013, p. 316). The ‘votes’ for a specific class can be interpreted as a measure of probability for that class occurring in a specific location. The final prediction is determined by the class with the highest probability (vote count) to occur in a specific location. The randomForest package (Liaw and Wiener, 2002) was used to perform the analysis.

RF generally performs well with default settings, i.e. without the tuning of parameters. Initial tuning of the number of trees in the forest (n_tree) and the number of variables to consider at any given split (mtry) showed a very limited impact on model performance, while at the same time the tuning process was very time-consuming. It was therefore decided to use the default parameter values.

The response variable is highly imbalanced (Table 2). This was accounted for by utilising a balanced version of RF (Chen et al., 2004). This is achieved by specifying the strata and sampsize arguments of the randomForest() function. The strata are the lithology classes and the sample size is determined by $c \cdot n_{min}$, where $c$ is the number of lithology classes (seven) and $n_{min}$ is the number of samples in the least frequent class. Hence, downsampling is applied when growing individual trees. However, each sample is drawn from all available observations as many trees are grown, making this scheme likely more effective than downsampling the dataset prior to model building.

RF also provides a relative estimate of predictor variable importance. The importance() function of the randomForest package allows to assess variable importance as the mean decrease in either accuracy or node purity. However, the latter approach might be biased when predictor variables vary in their scale of measurement or their number of categories (Strobl et al., 2007) and was not used here. Variable importance is therefore measured as the mean decrease in accuracy associated with each variable when it is assigned random but realistic values and the rest of the variables are left unchanged. The worse a model performs when a predictor is randomised, the more important that predictor is in predicting the response variable. The mean decrease in accuracy was left unscaled as recommended by Strobl and Zeileis (2008), and is reported as a fraction ranging from 0 to 1. Per default, individual trees of the forest are built using sampling with replacement (replace=TRUE). However, it has been shown that this choice might lead to bias in predictor variable importance measures (Strobl et al., 2007). It was therefore opted to use sampling without replacement.
3.5 Spatial cross-validation

Detailed guidelines for optimising sampling design for accuracy assessment have been developed (Olofsson et al., 2014; Stehman and Foody, 2019). However, this would require collecting new samples after modelling, which was not feasible given the geographic scope. Cross-validation schemes are frequently used to deal with such situations. It can be assumed that the response variable is spatially structured to some extent and cross-validation therefore requires accounting for the spatial structure (Roberts et al., 2017). Here, a spatial leave-one-out cross-validation (S-LOO CV) scheme was applied. In a conventional LOO CV, a single observation is removed from the dataset and all other observations (n – 1) are used to train the model. The class of the withheld observation is then predicted using the n – 1 model. This is repeated for every observation in the dataset, producing observed and predicted classes at every location. In a S-LOO CV scheme, a buffer is placed around the withheld observation and training data from within this buffer are omitted from both model training and testing so that there are no training data proximal to the test. The S-LOO CV scheme used here was adapted from Misiuk et al. (2019). The buffer size was estimated with the spatialAutoRange() function of the blockCV package (Valavi et al., 2018).

3.6 Accuracy assessment

The accuracy of the model was assessed based on a confusion matrix that was derived by the S-LOO CV. Overall accuracy and the balanced error rate (BER) were used to evaluate the global accuracy of the model, while error of omission and error of commission were selected as class-specific metrics of accuracy. The overall accuracy gives the percentage of cases correctly allocated and is calculated by dividing the total number of correct allocations by the total number of samples (Congalton, 1991). The BER is the average of the error rate for each class (Luts et al., 2010). The error of omission is the number of incorrectly classified samples of one class divided by the total number of reference samples of that class. The error of commission is the number of incorrectly classified samples of one class divided by the total number of samples that were classified as that class (Story and Congalton, 1986). The overall accuracy, its 95% confidence intervals and a one-sided test to evaluate whether the overall accuracy was significantly higher than the no information rate (NIR) were calculated by applying the confusionMatrix() function of the caret package (Kuhn, 2008). The confidence interval is estimated using a binomial test. The NIR is taken to be the proportion of the most frequent class. Errors of omission and commission are not provided by the function but can be calculated from the confusion matrix. The BER was calculated with the BER() function of the package measures (Probst, 2018).

4 Results

4.1 Variable selection

The Boruta algorithm was run with maxRuns = 500 iterations and a p-value of 0.05, leaving no variables unresolved (i.e. tentative). All 38 predictor variables initially included in the model were deemed important according to the Boruta analysis
mates over the correlation coefficient $r$, a value of 0.5 was selected (Fig. 3) This selection ensured high model performance while at the same time minimising the number of predictor variables. Subsequent correlation analysis reduced the number of retained predictor variables to eight. These were bathymetry (MS_bathy_5m), distance to shore (MS_biogo5_dist_shore_5m), sea-surface temperature range (BO2_temprange_ss), sea-surface maximum primary productivity (BO2_ppmax_ss), sea-floor minimum temperature (BO2_tempmin_bdmean), sea-surface maximum salinity (BO2_salinitymax_ss), sea-surface salinity range (BO2_salinityrange_ss) and sea-surface minimum silicate (BO2_silicatemin_ss). The strongest correlation between remaining predictor variables (Fig. 4) was found between bathymetry and sea-floor minimum temperature ($r = 0.38$), sea-surface maximum salinity and sea-surface minimum silicate ($r = -0.36$) and bathymetry and distance to shore ($r = -0.33$). Maps of the selected predictor variables are shown in Fig. A1.

4.2 Environmental space

The environmental space (Fig. 5) is generally sampled adequately, although there is a tendency for an over-representation of shallower water depths and areas closer to land. Sea-surface temperature range, sea-surface maximum primary productivity, sea-floor minimum temperature, and sea-surface maximum salinity are all slightly biased towards higher values. Sea-surface salinity range and sea-surface minimum silicate are the environmental variables that are most closely represented by the samples.

4.3 Model accuracy

The confusion matrix based on the S-LOO CV is shown in Table 3. The overall accuracy of the model is 59.4 %, with 95 % confidence limits of 58.4 % and 60.3 %. This is significantly higher ($p < 2.2e-16$) than the NIR (50.3 %). The BER is 0.54. The two dominant classes, Calcareous sediment, and Clay have the lowest error of commission with 18.3 % and 33.1 %, respectively. Calcareous sediment is most frequently mis-classified as Clay and vice versa. All other classes have high errors of commission (>70 %). Errors of omission are slightly higher than those of commission for the frequently occurring lithologies Calcareous sediment, and Clay, while lower for the rare classes Diatom ooze, Lithogenous sediment, and Radiolarian ooze.

4.4 Spatial distribution of deep-sea sediments

Probability surfaces of individual sediment classes with verbal descriptions of likelihood (Mastrandrea et al., 2011) based on the estimated probabilities are displayed in Fig. 6. For any given pixel in the map, the final lithology class is that one with the highest probability. The probability of the most probable class might be interpreted as a spatially explicit measure of map confidence. The resulting maps of the spatial distribution of deep-sea sediments and their associated confidence are shown in Fig. 7. Calcareous sediment and Clay dominate throughout the Pacific, Atlantic and Indian Oceans, whereby Clay occupies the deep basins and Calcareous sediment is found in shallower parts of the ocean basins. In the Southern Ocean, seafloor sediments are arranged in a banded pattern around Antarctica, with Lithogenous sediment forming an inner ring closest to land.
An outer ring of siliceous oozes (Diatom ooze, and Radiolarian ooze) dominates in the Southern Ocean. The width of this “opal belt” (Lisitzin, 1971) varies and in places, most notably south of South America, it is discontinuous. Overall, map confidence varies between 0.21 and 1. It is generally lower in the vicinity of class boundaries and higher in the geographic centre of a class.

The sea-floor lithology map bears a notable resemblance with previously published hand-drawn maps (e.g. Berger, 1974). The general patterns are very similar, e.g. the distribution of Calcareous sediment, Clay, and Diatom ooze in the major ocean basins.

Patterns of Radiolarian ooze in the Indian Ocean resemble those in Thurman (1997: Fig. 5-22). In the Pacific Ocean, Radiolarian ooze is mapped widespread in the vicinity of the equator, although not in the form of a narrow band as frequently depicted in hand-drawn maps (Berger, 1974; Thurman, 1997).

Based on the predicted distribution of lithology classes, Calcareous sediments cover approximately 121 million km$^2$ of seabed below 500 m water depth, equivalent to 36.8 % of the total area (Table 4). Clays are the second most frequent lithology occupying 102 million km$^2$ (31.0 %). Diatom ooze, Lithogenous sediment, and Radiolarian ooze account for 8.5 %, 9.5 % and 14.2 % of deep-sea floor, respectively.

4.5 Predictor variable importance

The three most important predictor variables were sea-surface maximum salinity, bathymetry, and sea-floor maximum temperature with mean decreases in accuracy above 5 % (Fig. 8). These findings are similar to results from Dutkiewicz et al. (2016), who determined sea-surface salinity, sea-surface temperature and bathymetry as the most important controls on the distribution of deep-sea sediments. Sea-surface minimum silicate was of medium importance (4.3 % decrease in accuracy), while sea-surface temperature range, sea-surface maximum primary productivity, distance to shore and sea-surface salinity range were of lower importance (<3 % decrease in accuracy).

5 Limitations of the approach

This study utilised legacy sampling data to make predictions of the spatial distribution of seafloor lithologies in the deep-sea. This is the only viable approach as it is unrealistic to finance and execute a survey programme that samples the global ocean with adequate density within a reasonable timeframe. However, this approach also has some drawbacks:

The presented spatial predictions were based on forming relationships between lithology classes and environmental predictor variables. For such a task, it would be desirable to cover the range of values of each of the predictor variables used in the model (Minasny and McBratney, 2006). Although it was not possible to design a sampling survey, it became nevertheless obvious that the environmental space is reasonably well covered, presumably because of the relatively large number of observations, which was achievable as there was virtually no cost associated with “collecting” the samples. However, it might not always be the case that a large sample dataset leads to adequate coverage of the environmental space. In such a case, it might be desirable to draw a suitable sub-sample that approximates the distribution of the environmental variables.
Data originating from many cruises over long time periods are most likely heterogeneous, which might lead to increased uncertainty in the predictions. Sources of uncertainty might relate to sampling gear type, vintage and timing of sampling, representativeness of subsampling, analytical pre-treatment, inconsistency of classification standards and more (van Heteren and Van Lancker, 2015). However, Dutkiewicz et al. (2015) made efforts to homogenise the data. From a total number of more than 200,000 samples, they selected 14,400 based on strict quality-control criteria. Only surface and near-surface samples that were collected using coring, drilling, or grabbing methods were included. Furthermore, only samples whose descriptions could be verified using original cruise reports, cruise proceedings and core logs were retained. Their classification scheme is deliberately generalised in order to successfully depict the main types of sediments found in the global ocean and to overcome shortcomings of inconsistent, poorly defined and obsolete classification schemes and terminologies (Dutkiewicz et al., 2015). Additional uncertainty might be introduced through imprecise positioning of the samples, which might lead to incorrect relations between the response variable and the predictor variables. No metadata exist on the positioning accuracy or even the method of determining the position, which might give some clues on the error associated with the recorded positions. However, the chances that this shortcoming leads to significant problems when making associations between target and predictor variables are relatively low, as the chosen model resolution of 10 km is relatively coarse when compared with positioning accuracy.

The initial choice of predictor variables was informed by the current understanding of the controls on deep-sea sedimentation (Dutkiewicz et al., 2016; Seibold and Berger, 1996). Consequently, all selected predictor variables were deemed important (Fig. 2). The three most important predictor variables (Fig. 8) are also in good agreement with Dutkiewicz et al. (2016). However, the large errors of omission and especially commission for the rare lithologies Diatom ooze, Lithogenous sediment, and Radiolarian ooze might indicate that the environmental controls are less well represented for these sediment types. Lithogenous sediment comprises a wide range of grain-sizes (silt, sand, gravel and coarser) and proximity to land might be an insufficient predictor. In fact, distance to shore had the second lowest variable importance (Fig. 8).

Sedimentation rates in the deep-sea typically range on the order of 1 – 100 mm per 1000 yrs (Seibold, 1975). The sample depths in the dataset used here might have ranged from core top to a few dm. The lithologic signal might therefore be integrated over timescales of approximately 100 yrs to a few 100,000 yrs. The model hindcasts to derive the oceanographic predictors typically cover approximately 25 yrs, while bathymetry and distance to coast might be nearly constant since global sea-level rise ceased approximately 6,700 yrs ago (Lambeck et al., 2014). Hence, there likely exists a mismatch between the time intervals, although oceanographic variables might not have changed dramatically over much longer timescales than a few decades.
6 Potential usage

Despite the good agreement with previously published maps and a reasonable overall map accuracy of \( \approx 60\% \), there is large variation in the class-specific error as well as the spatial distribution of map confidence. It is therefore recommended to always consult the information on map confidence along with the map of seafloor lithologies. The probability surfaces of the seven lithologies might be used as input for spatial prediction and modelling, e.g. marine species distribution modelling on a global scale, which typically lacks information on seafloor sediments, although substrate type is assumed to be an important environmental predictor. Additionally, the presented data layers might be useful for the spatial prediction of sediment properties (e.g. carbonate and organic carbon content).

The categorical map might serve as a resource for education and teaching, provide context for research pertaining to the global seafloor, support marine planning, management and decision-making and underpin the design of marine protected areas globally. Additionally, the provided lithology map might be useful for survey planning, especially in conjunction with confidence information to target areas where a certain lithology is most likely to occur. Conversely, areas of low confidence could be targeted to further improve the accuracy of and confidence in the global map of deep-sea sediments.

7 Data availability

The input sample data (five major classes below 500 m water depth) and the presented model results (probability surfaces of the five lithologies, lithology map and associated confidence map) are archived at https://doi.org/10.1594/PANGAEA.911692 (Diesing, 2020).

8 Conclusions

Based on a homogenised dataset of seafloor lithology samples (Dutkiewicz et al., 2015) and global environmental predictor variables from Bio-ORACLE (Assis et al., 2018; Tyberghein et al., 2012) and MARSPEC (Sbrocco and Barber, 2013) it was possible to spatially predict the distribution of deep-sea sediments globally. The general understanding about the controls on deep-sea sedimentation helped building a spatial model that gives a good representation of the main lithologies Calcareous sediment, Clay, Diatom ooze, Lithogenous sediment, and Radiolarian ooze. Further improvements should be directed towards the controls on the distribution of rarer lithologies (Diatom ooze, Lithogenous sediment, and Radiolarian ooze).

Author contribution

MD designed the study, developed the model code, executed the analysis, and wrote the manuscript.
Competing interests

The author declares no competing interests.

Acknowledgements

Thanks to Karl Fabian (NGU) and Benjamin Misiuk (Memorial University of Newfoundland) for providing valuable feedback. The reviews from Everardo González Ávalos and an anonymous reviewer were extremely helpful and led to significant improvements in the methodology and the manuscript.

References


Diesing, M., Kröger, S., Parker, R., Jenkins, C., Mason, C. and Weston, K.: Predicting the standing stock of organic carbon in


R Core Team: R: A Language and Environment for Statistical Computing, 2018.


<table>
<thead>
<tr>
<th>Environmental variable</th>
<th>Statistics</th>
<th>Unit</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bathymetry</td>
<td>mean</td>
<td>m</td>
<td>Sbrocco and Barber (2013)</td>
</tr>
<tr>
<td>Distance to shore</td>
<td>mean</td>
<td>km</td>
<td>Sbrocco and Barber (2013)</td>
</tr>
<tr>
<td>Sea-surface temperature</td>
<td>mean, min, max, range</td>
<td>°C</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface salinity</td>
<td>mean, min, max, range</td>
<td>PSS</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface dissolved oxygen</td>
<td>mean, min, max, range</td>
<td>mol m⁻³</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface primary productivity</td>
<td>mean, min, max, range</td>
<td>g m⁻³ day⁻¹</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface iron concentration</td>
<td>mean, min, max, range</td>
<td>µmol m⁻³</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface nitrate concentration</td>
<td>mean, min, max, range</td>
<td>mol m⁻³</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface phosphate concentration</td>
<td>mean, min, max, range</td>
<td>mol m⁻³</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-surface silicate concentration</td>
<td>mean, min, max, range</td>
<td>mol m⁻³</td>
<td>Assis et al. (2018)</td>
</tr>
<tr>
<td>Sea-floor temperature</td>
<td>mean, min, max, range</td>
<td>°C</td>
<td>Assis et al. (2018)</td>
</tr>
</tbody>
</table>
Table 2: Seafloor lithology classes used in this study, their abbreviations, their relationships to classes in Dutkiewicz et al. (2015) and the number and percentage of samples. Not included are Ash and volcanic sand/gravel, Mixed calcareous-siliceous ooze, Siliceous mud, Sponge spicules and Shells and coral fragments of the original classification.

<table>
<thead>
<tr>
<th>Lithology class</th>
<th>Abbreviation</th>
<th>Relation to Dutkiewicz et al. (2015)</th>
<th>No. of observation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcareous</td>
<td>Calc.Sed</td>
<td>Calcareous ooze</td>
<td>5251 (50.3 %)</td>
</tr>
<tr>
<td>sediments</td>
<td></td>
<td>Fine-grained calcareous sediment</td>
<td></td>
</tr>
<tr>
<td>Clay</td>
<td>Clay</td>
<td>Clay</td>
<td>3714 (35.6 %)</td>
</tr>
<tr>
<td>Diatom ooze</td>
<td>Dia.Ooze</td>
<td>Diatom ooze</td>
<td>623 (6.0 %)</td>
</tr>
<tr>
<td>Lithogenous</td>
<td>Lith.Sed</td>
<td>Gravel and coarser</td>
<td>751 (7.2 %)</td>
</tr>
<tr>
<td>sediment</td>
<td></td>
<td>Sand</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Silt</td>
<td></td>
</tr>
<tr>
<td>Radiolarian ooze</td>
<td>Rad.Ooze</td>
<td>Radiolarian ooze</td>
<td>99 (0.9 %)</td>
</tr>
</tbody>
</table>
Table 3: Confusion matrix. Observed (reference) classes are in columns, predicted classes in rows.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Calc.Sed</td>
<td>3727</td>
<td>685</td>
<td>8</td>
<td>134</td>
<td>10</td>
<td>4564</td>
<td>0.183</td>
</tr>
<tr>
<td>Clay</td>
<td>773</td>
<td>2008</td>
<td>48</td>
<td>159</td>
<td>14</td>
<td>3002</td>
<td>0.331</td>
</tr>
<tr>
<td>Dia.Ooze</td>
<td>235</td>
<td>210</td>
<td>295</td>
<td>295</td>
<td>34</td>
<td>1069</td>
<td>0.724</td>
</tr>
<tr>
<td>Lith.Sed</td>
<td>273</td>
<td>434</td>
<td>112</td>
<td>127</td>
<td>3</td>
<td>949</td>
<td>0.866</td>
</tr>
<tr>
<td>Rad.Ooze</td>
<td>243</td>
<td>377</td>
<td>160</td>
<td>36</td>
<td>38</td>
<td>854</td>
<td>0.956</td>
</tr>
<tr>
<td>Column total</td>
<td>5251</td>
<td>3714</td>
<td>623</td>
<td>751</td>
<td>99</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Error of omission</td>
<td>0.290</td>
<td>0.459</td>
<td>0.526</td>
<td>0.831</td>
<td>0.616</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 4: Breakdown of areal coverage by lithology types in the global ocean below 500 m water depth.

<table>
<thead>
<tr>
<th>Lithology</th>
<th>Number of pixels</th>
<th>Area ($10^6$ km$^2$)</th>
<th>Area (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcareous sediment</td>
<td>1,211,063</td>
<td>121.106</td>
<td>36.80</td>
</tr>
<tr>
<td>Clay</td>
<td>1,019,160</td>
<td>101.916</td>
<td>30.97</td>
</tr>
<tr>
<td>Diatom ooze</td>
<td>279,955</td>
<td>27.996</td>
<td>8.51</td>
</tr>
<tr>
<td>Lithogenous sediment</td>
<td>312,668</td>
<td>31.267</td>
<td>9.50</td>
</tr>
<tr>
<td>Radiolarian ooze</td>
<td>467,775</td>
<td>46.778</td>
<td>14.22</td>
</tr>
<tr>
<td>Sum</td>
<td>3,290,621</td>
<td>329.062</td>
<td>100</td>
</tr>
</tbody>
</table>
Figure 1: Locations of samples used in this study based on data from Dutkiewicz et al. (2015). Land masses are derived from ESRI (2010). **Hillshade topography is derived from GEBCO (2015).**
Figure 2: Results of the Boruta variable selection process. All environmental predictor variables had an importance significantly higher than the shadow variables (shadowMin, ShadowMean and shadowMax).
Figure 3: Influence of r-value on the out of bag error of a random forest model with default parameters. The size of the circles indicates the number of selected predictor variables (Npreds).
Figure 4: Correlation plot showing the correlation coefficients of the selected predictor variables.
Figure 5: A visual check to what extent the samples cover the environmental space. Blue: Samples; Red: Environmental data.
Figure 6: Probability surfaces of the five predicted lithologies. The verbal likelihood scale is based on Mastrandrea et al. (2011). Land masses are derived from ESRI (2010).
Figure 7: a) Predicted lithology classes and b) associated confidence in the predictions. Land masses are derived from ESRI (2010).
Figure 8: Random Forest variable importance.
Figure A1: Plots of the selected scaled predictor variables.
Figure A2: Predicted lithology classes in the Southern Ocean. Land masses are derived from ESRI (2010).