Preprocessing approaches in machine-learning-based groundwater potential mapping: an application to the Koulikoro and Bamako regions, Mali

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Received: 10 May 2021 – Discussion started: 30 June 2021
Revised: 4 October 2021 – Accepted: 30 November 2021 – Published: 18 January 2022

Abstract. Groundwater is crucial for domestic supplies in the Sahel, where the strategic importance of aquifers will increase in the coming years due to climate change. Groundwater potential mapping is a valuable tool to underpin water management in the region and, hence, to improve drinking water access. This paper presents a machine learning method to map groundwater potential. This is illustrated through its application in two administrative regions of Mali. A set of explanatory variables for the presence of groundwater is developed first. Scaling methods (standardization, normalization, maximum absolute value and max–min scaling) are used to avoid the pitfalls associated with reclassification. Noisy, collinear and counterproductive variables are identified and excluded from the input dataset. A total of 20 machine learning classifiers are then trained and tested on a large borehole database (n = 3345) in order to find meaningful correlations between the presence or absence of groundwater and the explanatory variables. Maximum absolute value and standardization proved the most efficient scaling techniques, while tree-based algorithms (accuracy > 0.85) consistently outperformed other classifiers. The borehole flow rate data were then used to calibrate the results beyond standard machine learning metrics, thereby adding robustness to the predictions. The southern part of the study area presents the better groundwater prospect, which is consistent with the geological and climatic setting. Outcomes lead to three major conclusions: (1) picking the best performers out of a large number of machine learning classifiers is recommended as a good methodological practice, (2) standard machine learning metrics should be complemented with additional hydrogeological indicators whenever possible and (3) variable scaling contributes to minimize expert bias.

1 Introduction

Today, 2.5 billion people across the world depend on groundwater for domestic supply (Grönwall and Danert, 2020). Groundwater is crucial in arid regions, such as the Sahel, where aquifers provide a permanent source of good-quality water during the months of the year in which rainfall and surface water are absent (Llamas and Martínez-Santos, 2005; Díaz-Alcaide et al., 2017). In a context of climate change, with rainfall expected to decrease and droughts likely to become more intense (Arneth et al., 2019), groundwater resources will be increasingly relied upon. This could well be the case in rural Mali, where access to drinking water and sanitation is a concern. In 2017, only 68% of the rural population had “at least basic” drinking water access, while 24% relied on unimproved water sources like unprotected springs and wells (UNICEF/WHO, 2019).

Groundwater potential mapping (GPM) is recognized as a valuable tool to underpin the planning and exploration of groundwater resources (Elbeih, 2015). There is no universal consensus as to what groundwater potential means. Thus, GPM may consist of developing spatial estimates of ground-
water storage in a given region, measuring the probability of finding groundwater or predicting where the highest borehole yields may occur (Díaz-Alcaide and Martínez-Santos, 2019). Recent years have witnessed a growing interest in groundwater potential studies across Africa, largely as a result of the need to achieve Sustainable Development Goal 6. The majority of these studies rely on a combination of remote sensing, geographic information systems and geophysics (Delgado, 2018; Adeyeeye et al., 2019; Magaña et al., 2018; Mpofu et al., 2020; Owolabi et al., 2020; Saadi et al., 2021; Al-Djazouli et al., 2021), whereas others directly interpret the information from borehole databases (Díaz-Alcaide et al., 2017).

There are two main approaches to GPM, namely expert-based decision systems and machine learning methods. Expert-based techniques have been used for a long time (DEP, 1993) and include multi-influencing factor approaches (Magesh et al., 2012; Nasir et al., 2018; Martín-Loeches et al., 2018), analytical hierarchy processes (Mohammadi-Behzad et al., 2019; Al-Djazouli et al., 2021) and Dempster–Shafer models (Mogaji and Lim, 2018; Obeidavi et al., 2021). Other frequently used expert methods are weight of evidence and frequency ratio analysis (Falah and Zeinivand, 2019; Boughariou et al., 2021). Machine learning is comparatively newer. A major difference between machine learning and expert approaches is that machine learning classification uses the advantages of artificial intelligence to find intricate associations among explanatory variables that might otherwise pass unnoticed. Hence, machine learning is well suited to map complex spatially distributed variables such as groundwater occurrence. The GPM literature showcases a wide variety of supervised classification approaches. For instance, Al-Fugara et al. (2020) used mixed discriminant analysis to map spring potential in a watershed of Jordan, much like Odzemir (2011) mapped spring potential in a Turkish basin by means of a logistic regression method. Random forests have proved adept at mapping groundwater potential, both in mountain bedrock aquifers (Moghaddam et al., 2020) and large metasedimentary basins (Martínez-Santos and Renard, 2020). Other supervised classification methods used in the field of GPM include boosted regression trees (Naghibi et al., 2016), support vector machines (Naghibi et al., 2017b), neural networks (Lee et al., 2012; Panahi et al., 2020) and ensemble methods (Naghibi et al., 2017a; Martínez-Santos and Renard, 2020; Nguyen et al., 2020).

GPM is founded on the assumption that groundwater occurrence can be partially inferred from surface features. Some of the most frequently used explanatory variables in GPM studies include lithology, geological lineaments, landforms, topography, soil, land use/land cover, drainage and slope-related variables, rainfall, and vegetation indices (Jha et al., 2007). Supervised classification algorithms are trained to find the associations between these variables and known groundwater data. Once the algorithms yield accurate predictions, their findings are extrapolated to estimate groundwater potential across a given study area.

The majority of machine learning GPM studies face two major shortcomings. First, as the number of available boreholes to train and test the algorithms is usually “small” and the number of explanatory variables can be comparatively high, a crucial issue is how to reclassify explanatory variables to minimize noise and decrease the variability of each conditioning factor. Ultimately, categorical and continuous variables need to be reclassified either as integers or in intervals. As reclassification relies heavily on the operator, this implies that bias may be incorporated from the beginning (Martínez-Santos and Renard, 2020).

A second problem is that the outcomes of machine learning studies are almost invariably assessed by means of standard big data metrics such as precision, recall, and area under the receiver operating characteristic curve (Pradhan, 2013; Naghibi et al., 2016; Chen et al., 2019). While useful, these may be of limited value in cases where the input dataset consists solely of unambiguous examples. Furthermore, there are question marks as to whether these metrics are truly representative when developing spatially distributed estimates (Martínez-Santos et al., 2021a). In some instances, using ad hoc calibration elements such as complementary borehole information can contribute to a better interpretation of the results.

The objective of this research is to build on the existing literature by presenting two methodological additions to machine learning GPM. First, we explore different scaling methods to avoid the pitfalls associated with the reclassification of explanatory variables; more specifically, we compare several automated approaches to data scaling, including standardization, normalization, maximum absolute value and max–min optimization (Pedregosa et al., 2011). The second novelty has to do with the way that the outcomes of this research are evaluated. In this context, borehole flow rates are used as a means to complement standard machine learning metrics, thereby providing additional robustness to predictions. Our method is described in Fig. 1 and is demonstrated through its application to map groundwater potential across two regions of Mali. The geographical setting also represents a second problem is that the outcomes of machine learning studies are almost invariably assessed by means of standard big data metrics such as precision, recall, and area under the receiver operating characteristic curve (Pradhan, 2013; Naghibi et al., 2016; Chen et al., 2019). While useful, these may be of limited value in cases where the input dataset consists solely of unambiguous examples. Furthermore, there are question marks as to whether these metrics are truly representative when developing spatially distributed estimates (Martínez-Santos et al., 2021a). In some instances, using ad hoc calibration elements such as complementary borehole information can contribute to a better interpretation of the results.

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are relatively uniform from a spatial perspective (27 °C), although seasonal oscillations are observed. The coolest temperatures occur in the south in the month of January (19 °C), and the hottest temperatures occur in the north in May (41 °C). There is a clear rainfall gradient, from 300 mm/yr in the north to 1200 mm/yr in the south.

Figure 3 shows the major geological domains of the study area (BGS, 2021). The rocks that make up the Precambrian craton (south) are composed mainly of gneiss, schist and quartzite, representing metamorphosed volcanic–sedimentary sequences. The original sedimentary layers, which include shale, arkose, gravel and conglomerate, were intercalated with volcanic rocks, such as basalt, gabbro, dolerite, rhyolite and tuff. Further north, metasedimentary rocks of the Proterozoic age, predominantly low to medium grade metamorphosed sandstones, with varying amounts of mudstone and limestone, take up over 50% of the study area. Volcanic outcrops (basalts and gabbros) are located in the central sector and at the northern end. Sedimentary rocks (sandstone, limestone and shale) of the Cambrian–Carboniferous age and Cretaceous–Tertiary age occur in the northern third of the study area. Quaternary fluvial deposits associated with the Niger River are observed along the riverbed (Traore et al., 2018).

From a hydrogeological perspective, four major aquifer units can be distinguished (Traore et al., 2018). These include basement aquifers, aquifers linked to fractures and intergranular porosity of consolidated sedimentary rocks (Precambrian and Paleozoic), aquifers formed in intrusive volcanic rocks, and aquifers in unconsolidated sedimentary materials (Fig. 4).

Basement aquifers are mostly located towards the south of the Koulikoro region. These are characterized by a thick weathered mantle. The average thickness of the weathered formation over the basement in this region is 10–50 m. Groundwater flows preferentially through the weathered mantle. Within this mantle, the lower part is generally more transmissive due to the lower clay content, and the upper part is less permeable to flow but can still be important as a groundwater reservoir. Fractures increase reservoir permeability, although their storage capacity is typically low (Martín-Loeches et al., 2018). Borehole yields range from 4 to 6 m$^3$/h (Traore et al., 2018).

The Precambrian metasedimentary materials are located in the central part of the Koulikoro region. Metasediments are considered a mixed-permeability aquifer: low-permeability layers provide higher storage, whereas more fractured layers present higher permeability and lower storage. Mean aquifer thickness ranges from 30 to 50 m, and the average yield varies from 5 to 10 m$^3$/h; however, some boreholes exceed 100 m$^3$/h. In the north, the fractured Paleozoic rocks allow water to flow through the sandstone and limestone layers. Average borehole yields are around 6 m$^3$/h, and the fractured horizons are about 40–45 m thick. Finally, unconsolidated sedimentary materials are composed of shales and argillaceous sandstone interbedded with limestone. The average borehole yields around 7 m$^3$/h. The thickness of the saturated zone ranges from less than 100 to over 400 m (Traore et al., 2018).
Figure 2. Study area: the Koulikoro and Bamako regions located in south-western Mali (map data © 2015 Google).

Figure 3. Geological map showing the main units with outcrops in the study area (adapted from BGS, 2021).
2.2 Data

Borehole data were provided by the National Directorate of Water, Mali (DNH, 2010). The database contains information on 5387 boreholes (3772 successful and 1615 unsuccessful), distributed across 1605 human settlements. The smallest unit of data aggregation is the village level. Available information includes the total number of boreholes per settlement, the borehole success rate, the mean flow rate (m$^3$/h) and the number of boreholes within a given flow rate range (less than 5 m$^3$/h, between 5 and 10 m$^3$/h, and over 10 m$^3$/h). In most cases, there is also information on the mean borehole depth (m), mean static water table depth (m) and mean electric conductivity (µS/cm). Figure 5 shows that the most common yields range between 1.5 and 4.5 m$^3$/h, while the mean borehole depth ranges between 50 and 80 m. This can be assumed to be the thickness of the weathered formation over the fresh granite and is roughly consistent with studies conducted in Burkina Faso (Courtois et al., 2010). Water table depth ranges mostly between 5 and 15 m.

There is a considerable number of villages with a 100% success rate (530), many of which have a single borehole (452). This raises the question of whether villages with a small number of boreholes are statistically representative, particularly in cases where the mean yield is low. Moreover, it creates an imbalance in the input dataset. Therefore, villages with fewer than five boreholes were omitted from the sample.

2.2.1 Target variable

For the purpose of algorithm training, groundwater potential is defined as the likelihood of a drilled borehole being successful. Successful boreholes are those that yield sufficient water to justify the installation of a hand pump (>0.5 m$^3$/h) (Foster et al., 2006). Therefore, the target variable is binary and can be interpreted as the presence or absence of groundwater.

Villages where more than 50% of wells were known to be successful were labelled “positive”. The positive classification also applies to those villages with more than one high-yield borehole (>10 m$^3$/h). The others were labelled “negative”. The resulting input dataset consisted of 650 villages: 390 were labelled positive and 260 were labelled negative. This comprises information from 3345 boreholes: 2101 were successful and 1244 were unsuccessful.

2.2.2 Explanatory variables

Groundwater potential depends largely on groundwater recharge. In turn, recharge is influenced by five main factors (Kumar, 1997; Jyrkama et al., 2002), namely climate (e.g. precipitation, temperature and potential evapotranspiration), soils (e.g. texture, saturated hydraulic conductivity and moisture capacity), land cover (e.g. vegetation density and type), geomorphology (e.g. surface slope and drainage density) and hydrology (e.g. streamflow and water table depth).
Rainfall is the principal source of groundwater recharge. Recharge depends on the precipitation rate as well as on the surface and subsurface factors that will allow or prevent infiltration. Soil is important because its characteristics (permeability, grain shape, grain size and the void ratio) control percolation. Higher infiltration potential is associated with sandy and gravelly soils, whereas clayey and silty soils rank among the least favourable for recharge (Díaz-Alcaide and Martínez-Santos, 2019). Integration of land use and land cover is often used in groundwater potential mapping studies because land use changes, which are mostly induced by human activities, affect hydrological dynamics (Díaz-Alcaide and Martínez-Santos, 2019). For instance, croplands and forests, located in the southern part of the study area, could be associated with high groundwater potential because ploughing, root development and biological activity favour infiltration. Areas close to permanent water bodies also tend to correlate with a higher groundwater potential (Naghibi et al., 2017a). In contrast, urban settlements and wastelands are assumed to have low groundwater potential due to the presence of impervious surfaces as well as to the absence of moisture (Magesh et al., 2012). Geomorphology may be useful in identifying features that may be favourable for groundwater infiltration and storage (Díaz-Alcaide and Martínez-Santos, 2019). Alluvial fans, sand dunes, weathering mantles, floodplains and other accumulations of unconsolidated materials are generally recognized as the most interesting geomorphological features from a groundwater point of view (Venkateswaran and Ayyandurai, 2015). In contrast, landforms such as inselbergs, scarp sand ridges may be considered of little interest.

Hydrological factors such as drainage density or water table depth also play an important role in groundwater recharge. High drainage density means that runoff can be evacuated quickly; therefore, infiltration is less likely. (Magesh et al., 2012; Fashae et al., 2014). In addition, a high drainage density can be assimilated to a higher erosion potential. Meijerink (2007) shows that parallels can be found between drainage density and soil permeability in certain geological settings. Water table depth is useful for mapping water tables to determine the main recharge and discharge zones of an aquifer.

A total of 19 explanatory variables were selected from an extensive review of the literature on GPM (Díaz-Alcaide and Martínez-Santos, 2019). Explanatory variables (Table 1) include lithology (Fig. 2), landforms, land use, soil, expected thickness (Fig. 5), rainfall, water table depth, vegetation-related indices (the normalized difference vegetation index, NDVI, and the normalized difference water index, NDWI), slope curvature, slope, topographic wetness index, stream power index, drainage density, distance from channels, clay content and the clay mineral alteration ratio (Fig. 6). An additional layer with mean borehole flow rates per village was developed for the purpose of calibrating the results.

The QGIS 3.0 r.geomorphon plugin (Jasiewicz and Stepinski, 2013) was used to prepare the landform map. This approach uses a digital elevation model (DEM) for the classification and mapping of landform features based on the princi-
Table 1. Explanatory variables used in GPM. The scale/resolution, acquisition time and source of data for each factor are provided.

<table>
<thead>
<tr>
<th>Explanatory variables</th>
<th>Scale/resolution</th>
<th>Time period</th>
<th>Source of data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alteration band ratio</td>
<td>30 m</td>
<td>7–16 March 2020</td>
<td>Own elaboration from Landsat 8</td>
</tr>
<tr>
<td>Clay content</td>
<td>250 m</td>
<td>n/a</td>
<td>SoilGrids250m 2.0</td>
</tr>
<tr>
<td>Curvature</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
<tr>
<td>Saturated thickness</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM and borehole database</td>
</tr>
<tr>
<td>Water table depth</td>
<td>30 m</td>
<td>2010</td>
<td>Own elaboration from borehole database</td>
</tr>
<tr>
<td>Distance from channels</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
<tr>
<td>Geology</td>
<td>1 million : 5 million</td>
<td>n/a</td>
<td>British Geological Survey</td>
</tr>
<tr>
<td>Geomorphology</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
<tr>
<td>Land use</td>
<td>300 m</td>
<td>2009</td>
<td>ESA Climate Change Initiative</td>
</tr>
<tr>
<td>Soil</td>
<td>1 million : 3 million</td>
<td>n/a</td>
<td>European Soil Data Centre</td>
</tr>
<tr>
<td>Rainfall</td>
<td>0.5°</td>
<td>1950–2009</td>
<td>CRU TS 3.21 dataset (University of East Anglia)</td>
</tr>
<tr>
<td>Drainage density</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
<tr>
<td>Thickness matrix</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Derived from DEM and borehole database</td>
</tr>
<tr>
<td>Elevation (DEM)</td>
<td>30.53 m</td>
<td>23 September 2014</td>
<td>SRTM</td>
</tr>
<tr>
<td>NDVI</td>
<td>30 m</td>
<td>7–16 March 2020</td>
<td>Own elaboration from Landsat 8</td>
</tr>
<tr>
<td>NDWI</td>
<td>30 m</td>
<td>7–16 March 2020</td>
<td>Own elaboration from Landsat 8</td>
</tr>
<tr>
<td>Slope</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
<tr>
<td>SPI</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
<tr>
<td>TWI</td>
<td>30.53 m</td>
<td>n/a</td>
<td>Own elaboration from DEM</td>
</tr>
</tbody>
</table>

The abbreviations used in the table are as follows: DEM – digital elevation model; NDVI – normalized difference vegetation index; NDWI – normalized difference water index; SPI – stream power index; TWI – topographic wetness index; ESA – European Space Agency; CRU TS – Climatic Research Unit (CRU) Time Series (TS); SRTM – Shuttle Radar Topography Mission; n/a – not applicable.

ple of pattern recognition, rather than on differential geometry. By default, the r.geomorphon plugin classifies landforms into 10 different categories. Because some of them can be expected to play a similar role in the context of GPM, these were subsequently regrouped into four categories (Fig. 6a): flat areas, gentle slopes, steep slopes and ridges.

Soil descriptions (Fig. 6b) of the study area were obtained from the European Soil Data Centre (Dewitte et al., 2013). About 45% of the region is characterized by the presence of pisoplinthic Plinthosols, a type of soil with plinthite (Fe-rich), strongly cemented to indurated concretions or nodules, a humus-poor mixture of kaolinitic clay, and other products of strong weathering (IUSS Working Group, 2015). It usually changes irreversibly to a layer with hard concretions or nodules or to a hardpan on exposure to repeated wetting and drying. Hypoluvic Arenosols are present in 20% of the total surface and are characterized by deep sandy soils, which explains their generally high permeability. These are residual sandy soils following weathering of rocks, generally rich in quartz, in situ. Nearly 13% of the study area is characterized by petric Plinthosols, which share multiple features with pisoplinthic Plinthosols but which, unlike the latter, are arranged into continuous or fractured sheets of connected concretions or nodules and are strongly cemented to an indurated layer starting ≤ 100 cm from the soil surface. Lithic Lep- tosols (very thin soils on continuous rock that are extremely rich in coarse fragments with continuous rock from ≤ 10 cm from the soil surface), haplic Lixisols (higher clay content in the subsoil than in the topsoil, as a result of pedogenetic processes) and eutric Regosols (very weakly developed mineral soils in unconsolidated materials) constitute about 15% of the study area.

Land cover seems to present a clear association with the precipitation gradient (Fig. 6c). The southern part is characterized by open broadleaf deciduous forests (ESA, 2010). The central part is characterized by an alternation of shrublands, mosaics of cropland vegetation and rainfed cropland. West of Bamako, in the sparsely populated mountains, there...
are forests mixed with shrublands. The northern part of the study area is dominated by cropland mosaics; further north, the landscape is made up of open grasslands, sparse vegetation and bare areas.

Boreholes in the study area are often drilled until the unaltered bedrock is reached. As a result, borehole depth can be a suitable proxy for aquifer thickness (Fig. 6d). Because the borehole database includes static level measurements, an expected saturated thickness layer was computed by subtracting one measurement from the other (Fig. 6d).

Satellite monitoring does not penetrate deep into the ground, but it provides information about features that may be associated with shallow groundwater (Díaz-Alcaide and Martínez-Santos, 2019). This can be important in the case at hand, where the borehole database shows that the static level remains at around 5–15 m below the surface (Fig. 7b). Vegetation-related indices can be useful in this context, particularly when computed at the end of the dry season (Fig. 7c-d). Take, for instance, the normalized difference vegetation index (NDVI; Fig. 7c), which is an estimate of vegetation vigour. This index is derived from the response of vegetation to red and visible infrared wavelengths (Xie et al., 2008). Similarly, the normalized difference water index (NDWI; Fig. 7d) is used as a measure of the amount of water in the vegetation or soil moisture (Xu, 2006). Based on Landsat 8 products, the NDVI and the NDWI are computed as per Eqs. (1) and 2 respectively, where B3 represents the green band (0.53–0.59 µm), B4 is the red band (0.64–0.67 µm) and B5 is the near-infrared band (0.85–0.88 µm).

\[
\text{NDVI (Landsat 8)} = \frac{(B5 - B4)}{(B5 + B4)} \tag{1}
\]
\[
\text{NDWI (Landsat 8)} = \frac{(B3 - B5)}{(B3 + B5)} \tag{2}
\]

The digital elevation model (DEM) was obtained from the radar-based Shuttle Radar Topography Mission (SRTM), with a resolution of 1 arcsec (30 m). Topography is a relevant factor in groundwater distribution, storage and flow. Surface runoff and infiltration are also constrained by surface features (Elbeih, 2015). In this case, the DEM was used to develop the curvature (Fig. 7e), slope (Fig. 7f), topographic wetness index (TWI; Fig. 7g) and stream power index (SPI; Fig. 7h). The topographic wetness index (TWI) represents the ease with which water may accumulate at the surface (Beven and Kirkby, 1979). Similarly, the stream power index provides a measure of the erosive power of flowing water (Moore et al., 1991). Other features computed from topographic data include the landforms (Fig. 6a) and drainage-related layers (Fig. 7i-j).

The channels extracted from the DEM are used to develop the drainage density and distance from channels maps. Drainage density is computed as the total length of the streams per catchment unit area. Distance from channels was developed by extracting all major channels into a separate layer and developing 500 m, 1500 m and 2500 m buffers.

Clay content in the first few metres of the surface largely determines the percolation of water into the aquifer. Therefore, an additional clay content layer (g/kg) in the top 2 m of the terrain was considered (Poggio and de Sousa, 2020). This layer is obtained by state-of-the-art machine learning methods that use global soil profile information and covariate data to model the spatial distribution of soil properties around the world (Fig. 7k). This layer provides information about subsurface clay content. To complement the information on clay content on the subsurface, an additional layer has been developed by combining bands 6 (short-wave infrared 1) and 7 (short-wave infrared 2) of Landsat 8 (Ouirzif et al., 2019). This layer provides information on clay content on the surface and the relationship with infiltration (Fig. 7l).

2.3 Predictive mapping software

MLMapper v2.0 is an upgrade of the software developed by Martínez-Santos and Renard (2020). MLMapper is a QGIS3 plugin that allows for the development of predictive maps of a given target variable based on a series of explanatory variables (Fig. 1). It relies on a variety of supervised learning classifiers from the scikit-learn 0.24.1 toolbox (Pedregosa et al., 2011). Version 2.0 uses 19 different algorithms, including support vector machines (SVC), linear support vector machines (LVC), logistic regression (LRG), the decision tree classifier (DTC), the random forest classifier (RFC), K-neighbour classification (KNN), linear discriminant analysis (LDA), Gaussian naïve Bayes classification (NBA), multi-layer perceptron neural network (MLP), the AdaBoost classifier (ABC), quadratic discriminant analysis (QDA), gradient boosting classification (GBC), the Gaussian process classifier (GPC), the ridge classifier (RID), the stochastic gradient descent linear classifier (SGD), perceptron (PER), the passive-aggressive classifier (PASSI), the nu-support vector classifier (nuSVC) and the extra trees classifier (ETC).

2.4 Preprocessing of explanatory variables

Machine learning algorithms are designed to find meaningful associations between the explanatory variables and a given target variable. A preprocessing of explanatory variables is important to avoid bias. Take, for instance, the issue of ranges. Rainfall in the study area oscillates between 300 and 1200 mm/yr, whereas NDVI and NDWI values only range between −1 and +1. As some algorithms attribute weight to explanatory variables based on sheer magnitude, scaling is advisable to prevent the rainfall layer from having an undue bearing on the results. Furthermore, those algorithms that use gradient descent as an optimization technique (logistic regression and neural networks) require data to be scaled, as range differences lead to different step sizes in the gradient descent formula for each feature and, as a result, computation time increases. Algorithms such as K-neighbours and support vector machines are affected by different ranges be-
cause these classifiers rely on distances between data points to determine their similarity.

Scaling is advocated as an essential part of algorithm training, as each subsequent procedure depends on the choice of unit for each feature (Huang et al., 2015). Furthermore, scaling is expected to transform feature values based on a defined rule, so that all scaled features have the same degree of influence (Angelis and Stamelos, 2000).

In this context, the rationale behind using preprocessing approaches is to rely on raw data as much as possible, instead of reclassifying them into intervals generated statistically or by expert criteria. Thus, four scaling methods were used: standardization, maximal absolute scaler (MaxAbs), maximal–minimal scaler (MaxMin) and normalization (Pedregosa et al., 2011).

The standardized explanatory variables are obtained by removing the mean and scaling to unit variance (Zheng and Casari, 2018). Centring and scaling happen independently on each explanatory variable by computing the relevant statistics on the samples in the training set. The mean and standard deviation are then stored to be employed on later data using transform. This is obtained by means of Eq. (3), where \( \tilde{x} \) is the new value of the sample, \( x \) is the old value of the sample, \( u \) is the mean of the training samples and \( s \) is the standard deviation of the training samples:

\[
\tilde{x} = (x - \mu) / s.
\]  

The MaxAbs method transforms features by scaling each explanatory variable to a given range. This estimator scales and translates each feature individually so that it lies within the range given in the training set (between 0 and 1 in the case at hand). Thus, this scaling method is robust to small standard deviations in feature data (Pedregosa et al., 2011). MinMax scaling is performed as per Eq. (4):

\[
\tilde{x} = (x - \min(x)) \left( \frac{\max(x) - \min(x)}{1} \right).
\]  

The MaxAbsScaler works in a similar way, but the training data are scaled to fall within the \([-1, 1]\) range by dividing by the largest maximum value of each feature. Finally, under the normalizer procedure, each sample is rescaled regardless of other samples so that its norm ("L1", "L2" or "max") equals 1 (Pedregosa et al., 2011). The L1 norm attempts to minimize the sum of the absolute differences between the target value and the estimated values. In turn, the L2 norm uses least squares, whereas the L-max norm rescales by the maximum of the absolute values. The L2 norm is by far the most common and was adopted as the procedure of choice.

2.5 Supervised classification routine

MLMapper incorporates a series of routines to enhance the conventional training and testing process (Fig. 1); these include collinearity checks, random search parameter fitting and recursive feature elimination. A selection of the best al-

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**Figure 6.** Explanatory variables used to predict the GPM: (a) geomorphology, (b) land use (A.a: artificial areas; W.b: water bodies; M.c.v: mosaic cropland vegetation; R.c: rainfed cropland; B.a: bare areas; O.g: open grassland; M.v/c: mosaic vegetation/cropland; C./O.s: close to open shrubland; B.e or s.f. and M.f/s: broadleaf evergreen or semi-deciduous forest and mosaic forest/shrubland), (c) soil (Eu.Cam.: eutric Cambisols; Eu.Nit.: eutric Nitrisols; Eu.Reg.: eutric Regosols; Hap.Lix.: haplic Lixisols; Hap.Ver.: haplic Vertisols; Hyp.Are.: hypoluvic Arenosols; Lit.Lept.: lithic Leptosols; Pet.Pl.: petric Plinthosols; Pis.Pl.: pisoplinthic Plithosols; Und.Gl.: undifferentiated Gleysols; Und.Fl.: undifferentiated Fluvisols; Vet.Cam.: vertic Cambisols) and (d) expected thickness matrix.
Collinearity occurs when two or more variables are highly correlated. This can affect the performance of the classifiers by attributing extra weight to an input variable or by adding noise to the final outcomes. Interpretability can also be impaired because the regression coefficients of certain algorithms are not uniquely determined (Martínez-Santos et al., 2021a). MLMapper incorporates a collinearity analysis function to prevent collinearity from adversely affecting the results. Collinearity analysis is performed before running the algorithms. Pairwise correlation among explanatory variables is computed, and correlation coefficients are expressed in a range between $-1.0$ (inverse correlation) and $1.0$ (direct correlation). Highly correlated explanatory variables may be excluded from the algorithm training procedure.

Random search parameter fitting increases the accuracy of the predictions by identifying the best combination of those parameters that govern each algorithm. The random search cross-validation function needs an algorithm, a scoring metric to evaluate the performance of the different hyperparameters, and a dictionary with the hyperparameter names and values. In regard to the previous version of MLMapper,
which used grid search cross-validation, this provides additional flexibility and reduces computation time. A sensitivity analysis of the number of iterations was performed. The best compromise between the results and computation cost was obtained by fixing the number of iterations at 500. No significant improvement in scoring metrics was observed for higher values, but running times increased considerably.

Once the optimal parameter distributions have been selected, recursive feature elimination cross-validation (RFECV) is performed. RFECV is only available for those algorithms that use feature importance or coefficient weight attributes. The goal of RFECV is to identify the most important explanatory variables for each algorithm as well as to eliminate those features that either incorporate noise or detract from the accuracy of predictions.

Classifiers are trained first on the initial set of explanatory variables. The weight of each feature is obtained, either through a coefficient attribute or through a feature importance attribute. The least important explanatory variables are then pruned from the model. This procedure is recursively repeated on the pruned set until the optimal number of features to be selected is eventually reached (Pedregosa et al., 2011). Once RFECV is complete, algorithms are run again on the input dataset (excluding noise and counterproductive variables) and hyperparameters are re-optimized. This step yields important information for the purpose of interpreting the maps, as it provides the user with the relative weight of explanatory variables.

Each algorithm predicts groundwater potential throughout the study area, assigning a value of 1 to pixels with positive groundwater potential and 0 to pixels where groundwater is expected to be absent. Each algorithm operates differently and relies on a different combination of explanatory variables, which inevitably leads to discrepancies in the predictions. In order to analyse the degree of agreement between the classifiers, an ensemble map is developed by computing the arithmetic mean at the pixel scale of those algorithms exceeding 0.85 predictive accuracy. Green pixels mean that all of the best-performing algorithms agreed on a positive groundwater potential outcome (arithmetic mean of 1). Conversely, red zones represent those pixels where all of the best-performing algorithms agreed on a negative groundwater potential (arithmetic mean of 0). Intermediate colours represent various degrees of agreement among the algorithms.

2.6 Machine learning metrics for algorithm evaluation

Outcomes of machine learning studies are often evaluated on the grounds of accuracy (test score) and the area under the receiver operating characteristic curve (AUC). The test score is calculated as the number of successful predictions over the total number of attempts in the test dataset, thereby providing a direct measure of predictive accuracy. In turn, the receiver operating characteristic curve shows the performance of a classification model at all classification thresholds. The AUC is a probabilistic metric that assesses the degree to which algorithms can distinguish between classes. Test score and AUC rank algorithms on a scale of 0 to 1, with a higher score implying better performance. If the classes are balanced, a score of 0.5 on either metric suggests that the prediction is no better than a random estimate.

Because the ratio between positive and negative classes in the input dataset is approximately 3:2, the F1 metric was also used to analyse the quality of the predictions in each of the classes separately. The F1 score is the harmonic mean of precision and recall. Precision is calculated like the ratio of true positives over the sum of true and false positives. Therefore, precision shows the classifier’s ability not to label a sample that is negative as positive. In turn, recall is the ratio of true positives over the sum of true positives and false negatives. In other words, recall represents the ability of the classifier to find all positive samples.

3 Results and discussion

3.1 Collinearity analysis

Figure 8 presents the results of the collinearity analysis. Pairwise correlation coefficients are expressed as a colour palette ranging from $-1.0$ (inverse correlation) to 1.0 (direct correlation). There is no hard threshold as to what is an acceptable level of correlation between two variables, although the literature shows that values between 0.4 and 0.85 may be acceptable (Dormann et al., 2013). In this case, no strong correlations were found among explanatory variables except when appraising the expected saturated thickness versus the expected thickness matrix. This pair renders a correlation coefficient of 0.85, which is close to the upper acceptability threshold. Because the number of non-surface features in the explanatory dataset is limited, a closer look was taken at the importance of these two variables. RFECV reveals that both are relied upon by the best-performing algorithms. In addition, the thickness matrix has seldom been used in the literature. These reasons led to the decision of keeping both variables.

3.2 Model evaluation and scaling method selection

Table 2 shows the results of the best algorithms for each scaling method. The five best algorithms for each scaling method were selected, and the mean of each metric was computed to analyse performance. All scaling methods render a mean test score for the top five algorithms in excess of 0.85, which implies that the predictions are accurate in all cases. The standardization scaling method presents the highest value (0.867), followed by the MaxAbs method (0.866), the normalization method (0.860) and the MaxMin method (0.857). All algorithms render a higher F1 score for the positive outcome, which is the most common one in the input database. However, the F1 score for the negative class is always higher.
than 0.81. This indicates that the algorithms are capable of distinguishing positive and negative classes accurately.

AUC exceeds 0.90 in all cases except for the AdaBoost algorithm (0.898) and decision tree algorithm (0.893). This implies that all of the best-performing algorithms have a high probability of adequately predicting the target variable. When comparing the scaling methods, the standardized one presents the highest test score and F1 score for the negative class, while the MaxAbs method presents the second-best performance in terms of the test score, F1 score for the negative class and AUC metrics. Hyperparameter optimization by the random search procedure led to the scores in Table 3.

The results in Table 2 show that the best scaling methods based on the score obtained in machine learning metrics are the standardized and MaxAbs techniques. Therefore, the focus of ensuing analyses will be placed on these two methods.

Noticeably, the best-performing classifiers all belong in the family of tree-based algorithms. Decision tree (DTC), random forest (RFC), AdaBoost classifier (ABC), gradient boosting (GBC) and extra trees (ETC) consistently outperformed other classification methods throughout the process. While tree-based algorithms do not necessarily require feature scaling (Li et al., 2017), the outcomes of this experience show that different scaling methods lead to slightly different performances. The best test scores were obtained by the ETC algorithm (0.878 and 0.872 for the standardized and MaxAbs methods respectively), whereas the test scores of the other algorithms remained above 0.85. The F1 score for the positive groundwater potential is above 0.87 in all cases, whereas the F1 score for the negative potential decreases slightly (0.82 to 0.85). AUC scores oscillate between 0.89 and 0.91, which reinforces the idea that the best-performing classifiers are able to accurately predict the target.

By nature, tree-based algorithms can be expected to yield good results in groundwater potential studies. DTC is a popular technique, largely due to the ease with which the internal logic of the algorithm can be interpreted (Naghibi et al., 2016; Moghaddam et al., 2020). In turn, RFC is a combination of tree predictors where each tree depends on the values of a random vector sampled independently and with the same distribution for all trees in the forest (Breiman, 2001). RFC is generally one of the most powerful machine learning classifiers and typically renders useful outcomes in predictive mapping (Naghibi et al., 2017b; Martínez-Santos and Renard, 2020). The GBC algorithm is an ensemble method in which multiple weak classifier trees are used to build a strong classifier (Friedman, 2001). In the same way, the basic principle of ABC (Freund and Schapire, 1997) is to fit a sequence of weak learners into repeatedly modified versions of the data. The predictions of all of them are combined by a weighted majority vote to produce the final prediction. The data modifications at each iteration of the so-called boosting consist of applying weights to each of the training samples (Pedregosa et al., 2011). The literature shows that GBC
Table 2. Results of the top five algorithms for each scaling method. Refer to section 2.6 for the definition of each metric: test score is the optimized test score, an F1 score of 0 means that the F1 score is false, an F1 score of 1 means that the F1 score is true and AUC refers to the area under the curve. The mean of each metric has been calculated for the different reclassification methods to compare the results.

<table>
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<th>Test score mean</th>
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<th>F1 score 0 mean</th>
<th>F1 score 1 mean</th>
<th>F1 score 1 mean</th>
<th>AUC mean</th>
<th>AUC mean</th>
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</table>

and ABC have been successfully used as ensemble methods in the development of GPM maps (Nguyen et al., 2020a; Martínez-Santos and Renard, 2020). Finally, for the ETC algorithm (Geurts et al., 2006), randomization goes a step further in the way splits are computed. As in random forests, a random subset of candidate features is used, but instead of looking for the most discriminative thresholds, random thresholds are extracted for each candidate feature and the best of these randomly generated thresholds is chosen as the splitting rule. This usually allows the variance of the model to be reduced somewhat further (Pedregosa et al., 2011).

Other algorithm families have been successfully applied in GPM research. Support vector machines (Martínez-Santos and Renard, 2020), logistic regression (Ozdemir, 2011; Chen et al., 2018) and neural networks (Moghaddam et al., 2020; Panahi et al., 2020) have all been demonstrated to be efficient predictors of groundwater potential, even when pitched against tree-based algorithms. This suggests that differences in input databases and the choice of explanatory variables may constrain the applicability of each algorithm in each given context. Furthermore, the inherent complexity of supervised classification makes it difficult for users to understand the internal logic of the algorithms, which means it is frequently impossible to predict which one will perform best in a given situation. Therefore, incorporating various algorithm families in GPM is perceived as a suitable course of action in GPM studies.

3.3 Importance of explanatory variables

Naghibi and Pourghasemi (2015) indicate that the importance of explanatory variables in groundwater potential mapping is considerably influenced by the approach used in an investigation and the study area properties. Outcomes show that elevation, rainfall, geology and drainage density, among others, are the most important factors conditioning the groundwater potential, which is in agreement with results obtained by other authors in different geographical contexts (Ozdemir, 2011; Naghibi and Pourghasemi, 2015; Nguyen et al., 2020b).

A major advantage of incorporating recursive feature elimination is that it eliminates part of the expert bias associated with the choice of explanatory variables. In this case, the fact that all variables are used by at least two of the best-performing algorithms suggests that the initial choice of explanatory variables was appropriate. However, feature selection reveals clear differences among the classifiers. Under standardized scaling, RFC only required three explanatory variables to predict groundwater potential (precipitation, expected saturated thickness and elevation). ABC and GBC used 8 each, whereas ETC and DTC used 11 and 14 respec-
Table 3. Algorithm hyperparameters (description after Pedregosa et al., 2011) and adjusted outcomes. The hyperparameters utilized are as follows: max_depth – the maximum depth of the tree; max_features – the number of features to consider when looking for the best split; min_samples_leaf – the minimum number of samples required to be at a leaf node; min_samples_split – the minimum number of samples required to split an internal node; random_state – controls both the randomness of the bootstrapping of the samples used when building trees and the sampling of the features to be considered when looking for the best split at each node; n_estimators – the number of trees in the ensemble methods; Algorithm – method used by AdaBoost (discrete SAMME boosting algorithm or real SAMME.R boosting algorithm); learning_rate – shrinks the contribution of each classifier.

<table>
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<th>Hyperparameter</th>
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<table>
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Classifier outcomes were extrapolated to produce groundwater potential maps. Figure 10 shows the groundwater potential predictions rendered by each of the five best-performing algorithms under the two most effective scaling methods. Red areas are those in which the algorithms have found a combination of explanatory variables leading to a negative potential. In turn, green zones represent a positive groundwater potential. All maps show a gradient characterized by the predominance of positive areas in the south to a greater proportion of negative areas in the north. This appears to be related to rainfall patterns. Other major negative zones occur around the mountain outcrops of the south-west. Low potential areas also occur around the mountain outcrops of the south-west. The large green zone around the southern region corresponds to the weathering mantle of basement rocks. Groundwater in basement aquifers is most often found in weathered formations and piedmonts of the outcrops. Piedmonts may exhibit high GPM because these are essentially a mixture of weathered and transported materials (Martin-Loeches et al., 2018). This area presents smooth orography, which means that high GPM is mainly determined by the weathered mantle. Previous research by Diaz-Alcaide et al. (2017) reported a medium GPM for the southern part of the Koulikoro region. Discrepancies between this and our results likely stem from the fact that these authors used a commune-based approach, rather than physical variables.

The central part of the agreement map shows a high potential for both methods, except in the higher altitude areas. This region, consisting of consolidated metasedimentary materials, has an average aquifer thickness of 30–50 m (Traore et al., 2018). High yields are associated with the weathered mantle developed in the upper part instead of with the predominant lithology. This becomes evident in the metasedimentary outcrops located in the highlands. These present a low groundwater potential because fracturing facilitates rapid groundwater percolation into the plains, where it accumulates in the alteration zone. Areas near major rivers, such as the Niger, Sankarani and Bani, also have a high potential. This is attributed to the high permeability of alluvial sediments. The northern part of the study area, formed by consolidated and unconsolidated sedimentary materials, has a low groundwater potential. Although geological conditions are more favourable for groundwater due to the type of materials than in other areas of the region, groundwater potential is limited by low rainfall. This is demonstrated by the feature importance analysis, which shows that precipitation is one of
Figure 9. Feature importance calculated for the best-performing tree-based algorithms using the standardized and MaxAbs scaling methods. The sum of all variable weights equals one.
the two most important explanatory variables for groundwater occurrence.

Overall, maps are remarkably similar, although the MaxAbs scaling method renders a slightly greater proportion of negative potential areas. The agreement map (Fig. 11) allows for an analysis of discrepancies among the best-performing algorithms. The MaxAbs scaling method predicts a low groundwater potential (pixel value 0–0.2) for 41% of the study area, while about 40% is identified as a high groundwater potential (>0.8) and the remaining 19% consists of zones with moderate potential (0.2–0.8). The standardized scaling method renders a high groundwater potential for 42%, low groundwater potential for 38% and various degrees of moderate potential for 20% of the region.

Martínez-Santos et al. (2021a) show that standard machine learning metrics do not necessarily provide an unambiguous measure of classifier performance when predicting spatially distributed outcomes. These authors argue that case-specific indicators could provide additional insights into the results. In this case, an independent dataset of groundwater data (borehole yield) was used to cross-check machine learning predictions. Villages featuring more than five boreholes (n = 334) were relied upon for this purpose. These villages were classified into three groups based on the predicted groundwater potential (“low” <0.2, “moderate” 0.2 to 0.8 and “high” >0.8). Table 3 shows the mean borehole success rate, the mean borehole yield and the percentage of boreholes with a flow rate in excess of 10 m$^3$/h in each category. The average water table depth is also provided for reference.

A first conclusion that springs to mind is that about 60% of the villages with more than five boreholes fall within the high-groundwater-potential description, while a further 20% present a moderate groundwater potential. To a certain extent, this was to be expected, as the larger populations develop in areas where groundwater is available. Table 3 also shows groundwater potential outcomes to match borehole flow rates reasonably.

The mean borehole success rate in areas labelled as high groundwater potential is 75.9% for the MaxAbs scaling method and 73.7% for the standardized method. In contrast, the mean success rate for villages in the low-groundwater-potential category is 33.6% and 32.1% for the corresponding methods respectively. These outcomes suggest that both scaling methods and, in general, the machine learning approach are adept at identifying areas where boreholes are more likely to be successful.

Both the mean borehole yield and the percentage of high-yield boreholes (>10 m$^3$/h) provide interesting benchmarks for comparison. The mean borehole yield is consistent with map outcomes. Under the MaxAbs scaling method, the mean borehole yield is 1.39 m$^3$/h in the villages located within low-groundwater-potential areas and 3.92 m$^3$/h in areas where groundwater potential is predicted to be high. Furthermore, 76.1% of the high-yield boreholes fall within high-groundwater-potential areas, whereas just 6.6% are located in low-groundwater-potential areas. Results are similar for the standardized scaling method: the mean borehole yield is 1.44 m$^3$/h and 3.63 m$^3$/h for the low- and high-potential categories respectively, whereas 74.6% of the high-yield boreholes are located in villages within high-groundwater-potential zones.

On a final note, the literature features few examples of groundwater potential studies in the study area. Perhaps the only systematic precedent is that carried out by Díaz-Alcaide et al. (2017). These authors performed a national-scale assessment of groundwater potential for the Republic of Mali based on the same borehole database that has been used in this research. However, they aggregated their data at the commune scale, rather than the village scale, and relied on different classification principles. Thus, their outcomes are to be interpreted at a different spatial resolution. This notwithstanding, their groundwater potential assessment for the Bamako and Koulikoro regions is similar to that obtained by means of machine learning classification. The lower-groundwater-potential areas were identified in the north, and the higher-groundwater-potential ones were found mostly in the south. This attests to the capacity of machine learning approaches to realistically depict field conditions and, thus, to underpin groundwater development.

### 3.5 Limitations

Predictive groundwater mapping presents some uncertainties. These are typically associated with deficiencies in data quality (e.g. small sample sizes, missing covariates, and biased and missing data) as well as with errors in the structural nature and specifications of the model (Rahmati et al., 2015). In spatial modelling studies, the sample size has proven to be a significant factor affecting the predictive abilities of the models (Guisan et al., 2007). Moghaddam et al. (2020) analysed the influence of sample size on GPM, concluding that there is a significant decrease in AUC values when the sample size is 25% of the input dataset. In the present case, with 650 ground-truth points, the effect of sample size could be ruled out as per this standard.

Input data were made available by government officers after careful field evaluation. From this perspective, the database is considered to be of high quality. However, it is also true that key hydrogeological variables are missing. Groundwater potential was evaluated in binary form (positive/negative) because the borehole database provides little information in terms of borehole productivity. Spatially distributed estimates of aquifer transmissivity, storage coefficients and yield would allow for a multi-class assessment, which, in turn, would provide a more realistic zoning of groundwater potential. Moreover, the village-scale resolution presents some shortcomings. As per the database, all boreholes within a given village have the same coordinates. This makes it difficult to train the algorithms when, for instance, the village overlies a non-homogeneous hydrogeological set-
Figure 10. Mapping outcomes of the top five supervised classification algorithms for the two best-performing scaling methods. The top row shows the MaxAbs scaling method for the (a) AdaBoost classifier, (b) gradient boosting classifier, (c) random forest classifier, (d) decision tree classifier and (e) extra trees classifier. The bottom row shows the standardized scaling method for the (f) AdaBoost classifier, (g) gradient boosting classifier, (h) random forest classifier, (i) decision tree classifier and (j) extra trees classifier.

Table 4. Results of the analysis based on the agreement map categories for the MaxAbs and standardized scaling methods. The analysis was performed for villages with more than five wells. The average success rate, average flow rate, average flow rate without negative boreholes, percentage of wells with a yield higher than 10 m$^3$/h and the average water table depth are shown.

<table>
<thead>
<tr>
<th>Groundwater potential</th>
<th>Number of villages</th>
<th>Borehole success rate (%)</th>
<th>Mean borehole yield (m$^3$/h)</th>
<th>Mean borehole yield – only positive boreholes (m$^3$/h)</th>
<th>Percentage of high-yield boreholes (&gt;10 m$^3$/h)</th>
<th>Average water table depth (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MaxAbs scaling method</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>70</td>
<td>33.6</td>
<td>1.39</td>
<td>3.58</td>
<td>6.6 %</td>
<td>18.6</td>
</tr>
<tr>
<td>Moderate</td>
<td>69</td>
<td>50.5</td>
<td>2.23</td>
<td>4.36</td>
<td>17.3 %</td>
<td>15.1</td>
</tr>
<tr>
<td>High</td>
<td>195</td>
<td>75.9</td>
<td>3.92</td>
<td>5.10</td>
<td>76.1 %</td>
<td>12.1</td>
</tr>
<tr>
<td><strong>Standardized scaling method</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>65</td>
<td>32.1</td>
<td>1.44</td>
<td>3.96</td>
<td>6.6 %</td>
<td>19.3</td>
</tr>
<tr>
<td>Moderate</td>
<td>70</td>
<td>55.5</td>
<td>2.78</td>
<td>4.57</td>
<td>18.8 %</td>
<td>16.1</td>
</tr>
<tr>
<td>High</td>
<td>199</td>
<td>73.7</td>
<td>3.63</td>
<td>4.87</td>
<td>74.6 %</td>
<td>11.8</td>
</tr>
</tbody>
</table>

https://doi.org/10.5194/hess-26-221-2022
Figure 11. Mapping outcomes of the agreement map for the MaxAbs scaling method and standardized scaling method: (a) training points on the MaxAbs scaling method agreement map, (b) training points on the standardized scaling method agreement map, (c) testing points on the MaxAbs scaling method agreement map and (d) testing points on the standardized scaling method agreement map.
ting. More detailed knowledge on how the boreholes are distributed in space would be expected to enhance the outcomes.

Along the same lines, the resolution of certain explanatory variables is potentially problematic. Take, for instance, soil and lithology, which are only available at the regional scale and might constrain groundwater potential to an important extent. Gómez-Escalonilla et al. (2021) explain that inroads can be made in the use of a dynamic explanatory variable. Take, for instance, evapotranspiration or seasonal fluctuations in VV (vertical transmit, vertical receive) and VH (vertical transmit, horizontal receive) polarization intensity (backscattering coefficient) and VV-polarization coherence (interferometric correlation) from the Sentinel-1 time series, from which temporal descriptors are derived. Furthermore, Worthington (2015) shows that groundwater modelling in bedrock aquifers is complex because there is often a substantial flow through fractures, and the inter-connectivity and magnitude of these fractures are usually uncertain. In this context, geophysical techniques can provide useful information that improves the ability to predict GPM. However, the absence of geophysical information also limits our knowledge about subsurface structures.

The lack of interpretability is a major drawback of machine learning algorithms. Except for the simple decision tree, whose internal logic is straightforward, all classifiers work according to a complex internal architecture. Therefore, it is nearly impossible to understand the reasoning that leads most algorithms to a given conclusion beyond computing feature importance. This, in turn, results in extensive trial and error before optimal outcomes are achieved.

Overfitting can lead to spurious results in machine learning models. Overfitting occurs when the model does not generalize well with the training data (i.e. it tries to fit the training data perfectly at the risk of missing the underlying associations between the explanatory and target variables). To address this, techniques such as splitting the initial dataset into separate training and test subsets, cross-validation, regularization and ensembles may be used (Dietterich, 1995; Yeom et al., 2018). Case-specific indicators may also be used to appraise the results beyond standard machine learning metrics, thereby adding to the robustness of predictions (Martínez-Santos et al., 2021a, b). In the case at hand, this is achieved by comparing map outcomes with the limited available data on borehole yield.

4 Conclusions

Machine learning applications are gaining recognition as valuable tools to underpin groundwater management. The ease with which machine learning algorithms find complex associations within large datasets as well as their ability to develop accurate predictions open up a whole new dimension to the analysis of groundwater data. Artificial intelligence approaches may improve water access in areas such as the Sa-


Data availability. Data can only be made available with previous authorization from the Direction Generale de l’Hydraulique of Mali.

Author contributions. VGE was responsible for conceptualizing the study, undertaking the formal analysis, developing the methodology and software, and writing and preparing the original draft of the paper. PMS conceptualized the study, developed the methodology and software, acquired funding, and reviewed and edited the paper. MML conceptualized the study and reviewed and edited the paper.

Competing interests. The contact author has declared that neither they nor their co-authors have any competing interests.

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Acknowledgements. The first author received an FPI grant from the Ministry of Science and Innovation to develop his PhD within this project (grant no. PRE2019-090026). The second author received a Salvador de Madariaga grant (grant no. PRX18/00235) from Spain’s Ministry of Education, Culture and Sport to carry out a 3-month research stay at the Université de Neuchâtel, Switzerland, where the original version of the software used in this paper was developed. The authors thank the Direction Generale de l’Hydraulique of Mali for making its borehole database available.

Financial support. This research has been supported by the Ministerio de Ciencia, Innovación y Universidades (grant no. RTI2018-099394-B-100).

Review statement. This paper was edited by Marnik Vanclooster and reviewed by Fatma Trabelsi and one anonymous referee.

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