Predicting osteoarthritis in adults using statistical data mining and machine learning

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Abstract

Background: Osteoarthritis (OA) has traditionally been considered a disease of older adults (≥65 years old), but it may appear in younger adults. However, the risk factors for OA in younger adults need to be further evaluated.

Objectives: To develop a prediction model for identifying risk factors of OA in subjects aged 20–50 years and compare the performance of different machine learning models.

Methods: We included data from 52,512 participants of the National Health and Nutrition Examination Survey; of those, we analyzed only subjects aged 20–50 years (n = 19,133), with or without OA. The supervised machine learning model 'Deep PredictMed' based on logistic regression, deep neural network (DNN), and support vector machine was used for identifying demographic and personal characteristics that are associated with OA. Finally, we compared the performance of the different models.

Results: Being a female (p < 0.001), older age (p < 0.001), a smoker (p < 0.001), higher body mass index (p < 0.001), high blood pressure (p < 0.001), race/ethnicity (lowest risk among Mexican Americans, p = 0.01), and physical and mental limitations (p < 0.001) were associated with having OA. Best predictive performance yielded a 75% area under the receiver operating characteristic curve.

Conclusion: Sex (female), age (older), smoking (yes), body mass index (higher), blood pressure [high], race/ethnicity, and physical and mental limitations are risk factors for having OA in adults aged 20–50 years. The best predictive performance was achieved using DNN algorithms.

Keywords: arthritis, machine learning, osteoarthritis, statistical data mining

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Introduction

Osteoarthritis (OA) affects a great number of people, affecting the society in terms of health and financial costs. Although OA has traditionally been considered a disease of older age, it can also affect younger adults, having a profound impact on their psychosocial well-being and work capacity. Early signs of OA can be present up to two decades prior to formal diagnosis. Data from the Global Burden of Disease (GBD) study indicate a steady growth in disease burden due to OA since 1990 among people aged 15–49 years, with the greatest burden evident for females. The rate of primary knee replacement for people aged 20–49 years increased by 76% in the United States from 2001 to 2007, and the prevalence of knee OA has more than doubled since the mid-20th century.2

Younger populations with OA represent a new public health issue, given that these individuals will likely live with OA for a longer time than previous generations. Moreover, early OA detection in younger adults is important for public health
because they tend to be more active than older adults, and OA has a heavy impact on work, sport, and quality of life. Identifying risk factors of OA in early stages among younger adults may help prevent severe OA through medical treatments, physiotherapy, joint viscosupplementation, or surgical procedures (e.g. valgus knee osteotomy, Bernese pelvic osteotomy).

Although there are several studies concerning risk factors for OA, only recently have machine learning methodologies begun to be used in OA research, especially with the population under 50. Current guidelines are not well suited for diagnosing patients in the early stages of OA and do not identify patients for whom OA might progress rapidly. To improve current practices, a comprehensive patient-specific risk models need to be developed and tested. Approaches such as data mining and machine learning will aid in the development of such models.

Since general OA is suspected by clinical examination and confirmed by diagnostic imaging (mostly X-ray), many studies focus on detecting OA using machine learning, specifically Neural Nets, for image learning. However, recent studies show that using statistical data based on demographic and personal characteristics without any medical images to predict the occurrence of diverse forms of OA can have a significant impact on preventive medical care (e.g. specific occupational or physical therapy) and on better choosing subjects who would need X-rays to avoid unnecessary irradiation.

Datasets in medicine are becoming larger; to analyze these vast amounts of data, researchers need to look beyond traditional statistical methods. A supervised machine learning model named ‘PredictMed’ has been developed and validated to predict health conditions in patients with developmental disorders. In the current study, we applied an upgraded version named ‘Deep PredictMed’ based on deep learning, logistic regression (LR), and support vector machine (SVM) for identifying risk factors of OA in subjects aged 20–50 years from the National Health and Nutrition Examination Survey (NHANES–USA national health survey). Therefore, the objectives of this study were to identify risk factors for OA in younger adults using machine learning and compare the performance of different machine learning models.

**Methods**

**Subjects, predictors, and subject selection**

**Subjects.** Subjects were selected (Figure 1) from the NHANES cohort (https://www.cdc.gov/nchs/nhanes/index.htm) which included 52,512 Americans aged 20 years or older. The guidelines of the ‘Transparent Reporting of a Multivariable Prediction Model for Individual Prognosis or Diagnosis Statement’ (TRIPOD) were followed.

**Outcomes.** Possible OA risk factors/associated conditions were analyzed among 31 variables from the NHANES data (Table 1). They were selected based on clinical experience and literature on OA risk factors. The NHANES includes data on age, gender, race/ethnicity, family, poverty income ratio (PIR), education level, cigarette smoking, physical activity, and various medical conditions. Poverty status was defined by using the poverty income ratio (PIR), an index calculated by dividing family income by a poverty threshold specific to family size. PIR was estimated using NHANES guidelines and adjustment for family size, year, and state of residence. PIR was reported by three levels: ≤1.3 (low), >1.3–3.5 (middle), and >3.5 (high).

Participants were asked about physical activity and were classified as being physically active if they reported walking, cycling, and performing moderate or vigorous work or leisure/home activity. Body mass index (BMI) was calculated as weight in kilograms divided by height in meters squared. Participants were considered to have a normal weight if they had a BMI <25 kg/m², to be overweight if they had a BMI between 25 and 30 kg/m², or obese if their BMI was 30 kg/m². Diabetes was defined by reported physician diagnosis, fasting plasma glucose >126 mg/dl, 2-h oral glucose tolerance test >200 mg/dl, or glycohemoglobin >6.5%. Hypertension was defined by reported physician diagnosis or mean systolic blood pressure (of up to four measurements on two separate occasions) >140 mmHg. Other comorbid conditions included reported diagnosis of myocardial infarction (MI) and stroke, and presence of physical and mental limitations caused by long-term physical, mental, and emotional problems or illness. The Physical Functioning Questionnaire (PFQ) was filled in by trained interviewers, using the
Figure 1. The selection process for the study population (NHANES: National Health and Nutrition Examination Survey).

Table 1. List of dependent and independent variables selected from the National Health and Nutrition Examination Survey.

| Presence of osteoarthritis | Sex                        | Age                  | Poverty income ratio                                      |
|----------------------------|----------------------------|----------------------|----------------------------------------------------------|
| Race/ethnicity             | Diabetes                   | Body mass index      | Hypertension if >140 mmHg                                 |
| Smoking status             | History of stroke          | Physical and mental limitation | Walked or bicycled over the past 30 days                  |
| Require special healthcare equipment | Difficulty in stooping, kneeling | Difficulty walking up 10 steps | Difficulty sitting for long periods                      |
| Difficulty walking for a quarter mile | Difficulty reaching up over head | Difficulty in performing leisure activity at home | Difficulty in performing house chore                      |
| Difficulty in lifting or carrying | Difficulty in standing for long periods | Difficulty in walking between rooms on same floor | Difficulty in dressing themselves                        |
| Difficulty in standing up from armless chair | Difficulty in using fork, knife, drinking from cup | Difficulty in grasping/holding small objects | Difficulty getting in and out of bed                      |
| Difficulty in preparing meals | Difficulty in attending social event | Difficulty in going out to movies, events | Difficulty in managing money                             |
Computer-Assisted Personal Interview (CAPI) system.\textsuperscript{14}

\textbf{Subject selection.} Selected 19,133 participants younger than 50 years were divided into categories with or without OA. As reported in a previous study\textsuperscript{14} on demographic and personal characteristics from NHANES database, the dependent variable OA was ascertained based on the question, ‘Has a doctor or other health professional ever told you that you have arthritis?’ and participants were classified as having OA if they reported ‘Osteoarthritis’ to the question ‘Which type of arthritis?’

\textbf{Data analysis.} Feature reduction, removing features with low variance,\textsuperscript{15} was performed and compared with feature generation by principal component analysis (PCA).\textsuperscript{16} LR,\textsuperscript{17} deep neural network (DNN),\textsuperscript{18} and SVM\textsuperscript{19–21} were used separately for predicting OA.

\textbf{DNN model development.} The 19,133 participants’ dataset with 31 raw features was split into three groups: Training (76.5\%), Test (15\%), and Validation (8.5\%). Figure 2 shows prediction algorithm flow diagram using a DNN model with scaled PCA data (with 31 raw features) split into training, testing, and validation sets; data preprocessing with a scaler was used to convert categorical variables into continuous variables, and PCA was employed to generate new features. The DNN model was trained with scaled PCA variables to generate a trained DNN model; model predictions were evaluated by comparison with the true data labeled by clinicians in the test set. We divided the training and test data to avoid overlapping of participant data.

Model performance was evaluated based on the number of true positives ($TP$), true negatives ($TN$), false positives ($FP$), and false negatives ($FN$). $TP$ was the correctly predicted OA participants, and $TN$ was the correctly predicted non-OA participants. $FP$ and $FN$ were the incorrectly predicted OA and non-OA participants, respectively. The two-classed dataset we used in this study was imbalanced because there were more subjects without (17,835) than with OA (1298). To deal with this, we used three metrics for proper evaluation:\textsuperscript{20} accuracy (Acc), sensitivity (Sn), and specificity (Sp). They were calculated using the following equations:

\begin{align*}
    \text{Sn} &= \frac{TP}{TP+FN}, \\
    \text{Sp} &= \frac{TN}{TN+FP}, \\
    \text{Acc} &= \frac{(TP+TN)}{(TP+FN+FP+TN)}
\end{align*}

In addition, we evaluated the area under the receiver operating characteristic (ROC) curve (AUC); it evaluates an algorithm performance using one single value.

\textbf{Data preprocessing.} PCA is a data processing methodology for extracting relevant information from datasets. By preprocessing data features, PCA obtains new features in a compact way: new features are the projection of original features onto few dimensions of the data space (the principal components, retaining the maximum data variance), so finding structures that enable a better fitting of classification algorithms.\textsuperscript{16}

The dataset used in this study included categorical/binary and continuous variables. To achieve a better classification performance of the DNN classifier, we converted all 31 binary or categorical variables into 31 continuous variables using scikit-learn Scaler methods:\textsuperscript{3} StandardScaler, RobustScaler, and QuantileScaler. We also tried not scaled PCA. Figures 3 and 4 show the first and second principal components of: (a) not scaled PCA and (b) PCA with Quantile Transformer Scaler.

We also studied the use of Feature Reduction (as an alternative method to PCA) by removing
features with low variance using the scikit-learn method `feature_selection.VarianceThreshold`. Out of 31 variables, the following 7 showed higher variance than the threshold (set empirically at 0.8): sex, age, cigarette smoking, BMI and high blood pressure (HBP), race/ethnicity, and physical and mental limitation. Next, we used the reduced dataset of 19,133 patients with the seven significant variables as input data to the DNN (scaled data).

**DNN structure.** A feed-forward neural network trained with standard backpropagation was used in several different configurations of learning models. For each model, hyperparameters were adjusted, including the number of hidden layers, the number of neurons in each layer, the activation function, the optimization method, the learning rate, the batch size, and regularization techniques. The DNN with the best predictive performance had four hidden layers with 50, 30, 8, and 2 neurons in each layer, respectively. The last output layer, with two neurons, used a Sigmoid activation function. The optimization loss function had accuracy metrics and `sparse_categorical_crossentropy` loss function. Adam optimization was used. The DNN used the SeLU activation function in each layer and Regularization (0.001) in each layer.

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**Figure 3.** Two-dimensional plot of first and second principal components of not scaled principal component analysis.

**Figure 4.** Two-dimensional plot of first and second principal components of quantile scaled principal component analysis. The first and second principal components in Figures 3 and 4 are the two main dimensions of variation of unscaled (Figure 3) and scaled (Figure 4) data, showing separation of two classes of people (0/1: with or without osteoarthritis).
We took into account the unbalanced nature of the dataset using the technique of ‘class weighting’,25,26 hence giving different weights to both the majority and minority classes. Class weights were used to fix the issue of unbalanced classes in the dataset, making the minority classes more important. The technique was implemented using dedicated Keras functions for deep learning.18

Epoch size and batch size of the training setting were set to 3000 and 256 sets, respectively. All models were implemented using Keras18 with TensorFlow and several python scikit-learn methods.27

We also evaluated other two methods to increase DNN performance: Dropout28 and Batch normalization11 techniques. Batch normalization helps to reduce sensitivity to initial starting weights. Dropout is used to push the model to a better generalization, reducing overfitting. Both Dropout and Batch normalization did not lead to a significant improvement in DNN performance.

**Comparison with LR and support vector classifiers.**

We compared the predictive performance of the DNN model with the Logistic Regression Classifier (LRC) and Support Vector Classifier (SVC) models.

**LRC:** We performed LR to identify the presence of OA based on sex, age, cigarette smoking, BMI and HBP, race/ethnicity, and physical and mental limitations.

The LRC performs the regression on a training test and test set, and accuracy is calculated. Then this step is repeated on a new training test and test set obtained by reshuffling previous training and test set, and accuracy is re-calculated. The process is repeated 20 times and the 20 accuracy values are finally averaged to obtain the final accuracy value. The same process is used to calculate final average sensitivity and specificity.

The following procedure was implemented:

- Split the 19,133 participants into training (80%) and test (20%) sets and perform LR to predict the probability of each patient of the test set to have OA.
- Classify each patient having a probability $p >$ threshold to be OA-positive.

Calculate accuracy, sensitivity, and specificity of the prediction.

Randomly re-shuffle the training and test sets, re-perform regression, and re-calculate accuracy, sensitivity, and specificity.

After 20 re-shuffles, calculate average accuracy, sensitivity, and specificity.

The LR algorithm was based on General Linear Model from R-programming language.30

**SVC:** On the same dataset used for LRC (19,133 patients, seven variables, plus OA as target variable), we split the 19,133 patients into training (80%) and test (20%) sets. We used an SV-Clat as our classifier (scikit-learn – svm.SVC) with a set-up taking into account the unbalanced dataset. After fine-tuning of the SVC hyperparameters (using scikit-learn SVM grid search method), we used a ‘poly’ kernel of degree seven with a cost $C = 10,000$ and a gamma_value = 0.0001. We randomly re-shuffled the training and test sets 10 times, each time calculating accuracy, sensitivity, and specificity.

**Results**

The models identified female sex, older age, race/ethnicity (Mexican American < Other Hispanic < Non-Hispanic White < Non-Hispanic Black < Other Race), being a smoker, having HBP and high BMI, and physical and mental limitations as factors significantly ($p < 0.01$) associated with OA in adults aged 20–50 years. We found the variables associated with OA using the WALD z-test (Table 2).21 DNN outperformed LRC and SVC. DNN using both scaled data and PCA showed the best results in terms of accuracy, sensitivity, specificity, and ROC curve area, similar to previous studies on OA detection with deep learning.16 The DNN after hyper-parameter tuning had 71% accuracy, 68% sensitivity, 71% specificity, and 75% ROC curve area (AUC). The ROC curve for the predictive performance of DNN with scaled PCA is shown in Figure 5, together with the Confusion Matrix showing the values of TP, TN, FP, and FN found by the DNN. This preprocessing showed a worse predictive performance when compared with the scaled PCA. Even using PCA after feature reduction did not show any improvement in comparison with simple scaled PCA preprocessing. The Dropout28 and Batch normalization30 techniques showed no significant impact on the DNN
performance. Training Loss and Validation Loss curves were compared, showing similar decreasing trends (Figure 6), revealing a good fit of the DNN learning algorithm and showing no over- or underfitting.

Results obtained using different input features are shown in Table 3. We repeated the comparison among DNN, LR, and SVM algorithms using same PCA features for all of them (Table 4). The combination of DNN and PCA with quantile transformer scaler showed the best predictive performance (Figure 3).

Discussion

Early signs of OA can be present in those aged between 15 and 49 years, up to two decades prior to formal diagnosis.1,2 Although there are several studies looking at risk factors for OA,3–35 only recently have machine learning methodologies begun to be used in OA research, and to the best of our knowledge, no studies are applying predictive models in the population under 50 years old.

We applied ‘Deep PredictMed’ models based on deep learning, LR, and SVM for identifying risk factors of OA in subjects aged 20–50 years from a large US national survey. The DNN overall predictive performance was better than the LRC and SVC models. The sensitivity of the LRC was lower than the DNN (50% versus 68%) since DNN deals better with unbalanced data.

In accordance with previous studies5,36,37 with older patients, the present research confirmed age, female sex, obesity, ethnicity, and smoking as risk factors of OA in patients aged 20–50 years. As reported in current literature, we also found that early OA was associated with increasing age and female sex.1–4,31,32 The increase in OA with age is a consequence of cumulative exposure to risk factors and biological changes such as oxidative damage, thinning of cartilage, or muscle weakness.33

There are known racial/ethnic differences in radiographic OA features. Our study confirms that

Table 2. List of the logistic regression coefficients (independent variables) associated with osteoarthritis.

| Independent variables | Logarithmic regression | Coefficient | Odds ratio | Standard error | Z ratio | Prob (>|z|) | p-value |
|-----------------------|------------------------|-------------|------------|----------------|---------|-------------|---------|
| 1. Intercept          |                        | -7.65       | 0.01       | 0.23           | -33.25  | <2.2e-16    | <0.001  |
| Female gender         |                        | 0.43        | 1.54       | 0.06           | 7.16    | 8.049e-13  | <0.001  |
| Age                   |                        | 0.08        | 1.09       | 0.01           | 21.31   | <2.2e-16    | <0.001  |
| Race/ethnicity        |                        | 0.06        | 1.06       | 0.02           | 2.56    | 0.01043     | 0.01    |
| Poverty income ratio  |                        | -0.01       | 1.00       | 0.01           | -0.65   | 0.51325     | .5      |
| Body mass index       |                        | 0.04        | 1.05       | 0.01           | 10.16   | <2.2e-16    | <0.001  |
| Cigarette smoking     |                        | 0.19        | 1.21       | 0.03           | 5.51    | 3.476e-08   | <0.001  |
| High blood pressure   |                        | 0.24        | 1.28       | 0.05           | 4.71    | 2.461e-06   | <0.001  |
| Physical and mental limitation | | -0.03 | 0.97 | 0.01 | -3.97 | 6.914e-05 | <0.001 |

**Logistic regression:** Female gender, being smoker, type of race/ethnicity (Mexican American < Other Hispanic < Non-Hispanic White < Non-Hispanic Black < Other race), the increasing of age, body mass index, and high blood pressure (positive values), and decreasing of poverty income ratio and physical and mental limitation (negative values) are predictive factors of osteoarthritis (in the ‘Estimate’ column). As an example, this means that for every unit increase in the female gender, the log odds \(\ln(p/1-p)\) increases 1.54 times (where \(p\) = probability to develop osteoarthritis), while for every unit decrease in poverty income ratio, the log odds \(\ln(p/1-p)\) decreases 1.00 time. The ‘Pr(>|z|)’ column at the far right in the table indicates the significant strength of the respective parameter in terms of \(p\)-value as osteoarthritis predictor. This means that the significance of female gender, age, race/ethnicity, body mass index, cigarette smoking, high blood pressure, and physical and mental limitation in predicting osteoarthritis is very probable, with a \(p\)-value <0.05.
African Americans are more likely to develop OA. Moreover, we found that compared with other ethnic groups, Mexican Americans and other Hispanics are less likely than young non-Hispanic Blacks and non-Hispanic Whites to have early OA. We also confirmed that HBP, cigarette smoking, and high BMI are associated with early onset of OA in younger adults. This highlights the need for preventive measures concerning general health (smoking cessation, improved nutrition, and physical activity) for preventing OA in younger adults. This also highlights the correlation between good physical condition and delay of the onset of OA. Another interesting finding was the correlation between early OA and physical and mental limitations, to be investigated more in depth in the future. However, from our results, we cannot exclude a reverse causal mechanism, that is, the physical and mental limitation should increase the risk of OA or vice versa.

Data mining and artificial neural networks can help the clinical decision-making process and precision medicine in the prediction of OA in people less than 50 years old. The proposed method identifies risk factors of having OA with indirect or limited data, such as the statistical data of medical utilization and health behavior information. This can be advantageous for possible patients to prevent future medical costs, reduce the time for diagnosis, and avoid unnecessary testing. Of course, no method is a panacea, and analysis of large datasets can also generate seemingly meaningful results which are in fact spurious artifacts. As datasets become ever larger and incorporate more complex variables, it becomes

| Table 3. Metrics comparison of LRC, DNN, and SVM algorithms (different input features). |
|-----------------------------------------------|
| Accuracy (%) | Sensitivity (%) | Specificity (%) |
| LRC          | 78             | 50             | 80             |
| SVM          | 56             | 80             | 55             |
| DNN          | 71             | 68             | 71             |

DNN, deep neural network; LRC, logistic regression classifier; SVM, support vector machine.

| Table 4. Metrics comparison of LRC, DNN, and SVM algorithms (same PCA input features). |
|-----------------------------------------------|
| Accuracy (%) | Sensitivity (%) | Specificity (%) |
| LRC          | 67.6           | 69.3           | 67.4           |
| SVM          | 62.3           | 68.9           | 61.8           |
| DNN          | 71             | 68             | 71             |

DNN, deep neural network; LRC, logistic regression classifier; SVM, support vector machine.
increasingly important to link confirmatory analysis with the scientific discovery process while incorporating study design and subject area expertise.\textsuperscript{4}

PredicitMed algorithm can help identifying subjects at risk and, therefore, anticipating X-ray and diagnosis of OA to propose conservative treatments instead of joint replacement. These results from the machine learning methods allow for the prediction of the presence of OA with good accuracy. The first clinical applications should be as follows:\textsuperscript{43,44}

In the case of osteo-articular pain and/or stiffness, the general practitioner (GP) could be alerted if the subject presents one or more risk factors, in which case medical imaging could be quickly proposed to confirm OA, and if present, specific treatment.

A score should be developed to help GPs, occupational physicians, and sports doctors better assess the risk of early OA to improve counseling and orientation of the type of work and sport activities.

Global health prevention measures concerning obesity, smoking, and hypertension are confirmed as important for early OA prevention, beyond their well-known role in cardiac and general health.

Early OA can be treated with non-surgical means such as viscosupplementation and medications (chondroitin sulfate, etc.); moreover, some localizations of early OA (e.g. knee and hip) are available for joint preservation through conservative surgical treatments such as osteotomies, whereas late discovery usually results in joint resection and arthroplasty.\textsuperscript{34,35}

\textbf{Limitations and future directions}

Despite similar studies on OA\textsuperscript{14,33} using NHANES database have been conducted, survey data have limitations. Self-reported data may lack objectivity. For instance, a subject could have assumed ‘osteoarthritis’ even if the diagnosis is different; on the other hand, a subject could have real symptomatic OA and ignore it. Moreover, this analysis does not take into account the various localizations of OA but only its presence or absence, whereby we planned to investigate the type, the topography, and the progression of OA and knee OA on the Osteoarthritis Initiative database and on a prospective cohort in a future study.

Furthermore, only available variables were explored. Many participants were excluded from the dataset due to unavailable data. Most input features were of binary type. Even if scaled PCA were used to improve data separation (Figures 3 and 4), new additional feature inputs would be required for optimization. In addition, we excluded subjects who were receiving treatment for OA. This issue may reduce the overall accuracy of the prediction model. Another limitation is that we cannot attribute cause–effect relationships. From our study, it is unclear whether physical or mental disability is a risk factor or a consequence of OA. We plan to develop this study in the future.

After developing a prediction model, external validation is strongly recommended to evaluate the performance of the model in other participant data.\textsuperscript{13,45} Such external validation requires that, for each individual in the new dataset, outcome predictions be made using the original model and be compared with the observed outcomes.\textsuperscript{45,46} We plan to validate the present model in a future study with the Osteoarthritis Initiative dataset.

The next steps of our study will also include three main goals: improve the quality of the data, the predictive performance of the model, and the prediction of knee OA progression. Regarding the quality of the data, we aspire to obtain increasingly selected clinical features associated with OA to define more precisely the OA patient type and to avoid missing data.

The DNN predictive model could also be trained (and tested) on different datasets having different features; we can build these datasets based on our specific research goals. For example, we could use as dataset features age, BMI, and gender only (or other similar combinations of features) and we could use these data to train and test the model, verifying the predictive performance of the model itself.

To improve the predictive performance of the model, we plan to replace the DNN with an Ensemble of classifiers (like a stacking of Neural Nets or a combination of DNN with SVM and/or LR). In addition to this, we also plan to study how to implement our models (DNN, SVM, or Ensemble) to other datasets of patients with
diabetes. Another possible development would be to test this model on a set of non-American subjects and using the same training and test set splits for all the models.

Based on the DNN-PredictMed model presented in this study, in future we aim to develop a predictive algorithm that, after a training phase, having as input the patient data like age, gender, presence of associated pathologies, weight, and so on will be able to provide as output the probability for that patient to develop OA in a given time-frame (3–5 years) after patient data have been taken. We think that this kind of algorithm, if implemented in a user-friendly web application (or a similar tool), could be of great help in supporting clinical decision based on a more accurate prediction of OA developments.

**Conclusion**
Gender (female), age (older), smoking (yes), BMI (higher), blood pressure (high), race/ethnicity (Mexican American < Other Hispanic < Non-Hispanic White < Non-Hispanic Black < Other Race), and physical and mental limitations are risk factors for OA in adults aged 20–50 years. The best predictive performance was achieved using DNN algorithms.

**Ethics approval and consent to participate**
This study did not require an ethical board approval because of involving information freely available in the public domain [National Health and Nutrition Examination Survey (NHANES)—USA national health survey]. The data were anonymized and informed written informed consent was obtained at the time of original data collection at participating sites.

**Consent for publication**
Not applicable.

**Author contributions**
**Carlo M. Bertoncelli**: Conceptualization; Data curation; Methodology; Writing – original draft.

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**Availability of data and materials**
Data available on https://www.cdc.gov/nchs/nhanes/index.htm

**References**
1. Ackerman IN, Kemp JL, Crossley KM, et al. Hip and knee osteoarthritis affects younger people, too. *J Orthop Sports Phys Ther* 2017; 47: 67–79.
2. Kurtz SM, Lau E, Ong K, et al. Future young patient demand for primary and revision joint replacement: national projections from 2010 to 2030. *Clin Orthop Relat Res* 2009; 467: 2606–2612.
3. Jamshidi A, Pelletier JP and Martel-Pelletier J. Machine-learning-based patient-specific prediction models for knee osteoarthritis. *Nat Rev Rheumatol* 2019; 15: 49–60.
4. Deveza LA, Nelson AE and Loeser RF. Phenotypes of osteoarthritis: current state and future implications. *Clin Exp Rheumatol* 2019; 37: 64–72.
5. Lim J, Kim J and Cheon S. A deep neural network-based method for early detection of osteoarthritis using statistical data. Int J Environ Res Publ Health 2019; 16: 1281.

6. Rajkomar A, Dean J and Kohane I. Machine learning in medicine. N Engl J Med 2019; 380: 1347–1358.

7. Bertoncelli CM, Altamura P, Vieira ER, et al. PredictMed: a logistic regression-based model to predict health conditions in cerebral palsy. Health Inform J 2020; 26: 2105–2118.

8. Bertoncelli CM, Solla F, Loughenbury PR, et al. Risk factors for developing scoliosis in cerebral palsy: a cross sectional descriptive study. J Child Neurol 2017; 32: 657–662.

9. Bertoncelli CM, Bertoncelli D, Elbaum L, et al. Validation of a clinical prediction model for the development of neuromuscular scoliosis. Pediatr Neurol 2018; 79: 14–20.

10. Bertoncelli CM, Altamura P, Vieira ER, et al. Predictive model for gastrostomy placement in adolescents with developmental disabilities and cerebral palsy. Nutr Clin Pract 2019; 35: 201–210.

11. Bertoncelli CM, Bertoncelli D, Maidan LM, et al. Identifying factors associated with intellectual disabilities in teenagers with cerebral palsy using a predictive learning model. J Child Neurol 2019; 34: 221–229.

12. Bertoncelli CM, Altamura P, Vieira ER, et al. Using artificial intelligence to identify factors associated with autism spectrum disorder in adolescents with cerebral palsy. Neuropediatrics 2019; 50: 178–187.

13. Moons KG, Altman DG, Reitsma JB, et al. New guideline for the reporting of studies developing, validating, or updating a multivariable clinical prediction model: the TRIPOD statement. Adv Anat Pathol 2015; 22: 303–305.

14. Mendy A, Park J and Vieira ER. Osteoarthritis and risk of mortality in the USA: a population-based cohort study. Int J Epidemiol 2018; 47: 1821–1829.

15. Girish C and Sahin F. A survey on feature selection methods. Comput Electr Eng 2014; 40: 16–28.

16. Smith LI. A tutorial on principal components analysis. Computer science technical report no. OUCS-2002-12, 2002, http://www.cs.otago.ac.nz/cosc453/student_tutorials/principal_components.pdf

17. Kleinbaum DG and Klein M. Logistic regression. New York: Springer, 2002.

18. Gulli A and Sujit P. Deep learning with Keras. Birmingham: Packt Publishing, 2017.

19. Amarappa S and Sathyanarayana SV. Data classification using support vector machine (SVM), a simplified approach. Int J Electron Comput Sci Eng 2014; 3: 435–445.

20. Baratloo A, Hosseini M, Negida A, et al. Part 1: simple definition and calculation of accuracy, sensitivity and specificity. Emerg 2015; 3: 48–49.

21. Gourieroux C, Holly A and Monfort A. Likelihood ratio test, Wald test, and Kuhn-Tucker test in linear models with inequality constraints on the regression parameters. Econ J Econ Soc 1982; 50: 63–80.

22. Kingma DP and Ba J. Adam: a method for stochastic optimization. In: Proceedings of the 3rd international conference for learning representations, San Diego, CA, 2015, https://arxiv.org/abs/1412.6980

23. Alcantara G. Empirical analysis of non-linear activation functions for Deep Neural Networks in classification tasks. arXiv preprint, arXiv:1710.11272, 2017, https://arxiv.org/abs/1710.11272

24. Michelucci U. Regularization. In: Michelucci U (ed.) Applied deep learning. Berkeley, CA: Apress, 2018, pp. 185–216.

25. Byrd Z. What is the effect of importance weighting in deep learning. Int Conf Mach Learn 2019; 97: 872–881.

26. Fu X. Training RBF neural networks on unbalanced data. In: Proceedings of the 9th international conference on neural information processing, vol. 2. New York: IEEE, 2002, https://ieeexplore.ieee.org/abstract/document/1198214

27. SciKit-Learn. A Python library used for data processing, https://scikit-learn.org/stable/

28. Jindal I, Nokleby M and Chen X. Learning deep networks from noisy labels with dropout regularization. In: Proceedings of the 2016 IEEE 16th international conference on data mining (ICDM), Barcelona, 12–15 December 2016. New York: IEEE.

29. Chambers J. Software for data analysis: programming with R. Cham: Springer, 2008.

30. Ioffe S and Szegedy C. Batch normalization: accelerating deep network training by reducing internal covariate shift. arXiv preprint, arXiv:1502.03167, 2015, https://arxiv.org/abs/1502.03167

31. National Institute for Health and Care Excellence. Osteoarthritis: care and management.
in adults. Clinical guideline CG177. London: National Institute for Health and Care Excellence, 2014.

32. Cross M, Smith E, Hoy D, et al. The global burden of rheumatoid arthritis: estimates from the Global Burden of Disease 2010 study. *Ann Rheum Dis* 2014; 73: 1316–1322.

33. Park J, Mendy A and Vieira ER. Various types of arthritis in the United States: prevalence and age-related trends from 1999 to 2014. *Am J Public Health* 2018; 108: 256–258.

34. Keenan OJF, Holland G, Maempel JF, et al. Correlations between radiological classification systems and confirmed cartilage loss in severe knee osteoarthritis. *Bone Joint J* 2020; 102-B: 301–309.

35. De l Escalopier N, Anract P and Biau D. Surgical treatments for osteoarthritis. *Ann Phys Rehabil Med* 2016; 9: 227–233.

36. Litwic A, Edwards MH, Dennison EM, et al. Epidemiology and burden of osteoarthritis. *Br Med Bull* 2013; 105: 185–199.

37. Blagojevic M, Jinks C, Jeffery A, et al. Risk factors for onset of osteoarthritis of the knee in older adults: a systematic review and meta-analysis. *Osteoarthritis Cartilage* 2010; 18: 24–33.

38. Nelson AE, Braga L, Renner JB, et al. Characterization of individual radiographic features of hip osteoarthritis in African American and White women and men: the Johnston County Osteoarthritis Project. *Arthritis Care Res* 2010; 62: 190–197.

39. Allen KD and Golightly YM. State of the evidence. *Curr Opin Rheumatol* 2015; 27: 276–283.

40. Zhang Y and Jordan JM. Epidemiology of osteoarthritis. *Clin Geriatr Med* 2010; 26: 355–369.

41. Vina ER and Kwoh CK. Epidemiology of osteoarthritis: literature update. *Curr Opin Rheumatol* 2018; 30: 160–167.

42. Collins GS, Reitsma JB, Altman DG, et al. Transparent reporting of a multivariable prediction model for individual prognosis or diagnosis (TRIPOD): the TRIPOD statement. *BMJ* 2014; 350: g7594.

43. Tognolo L, Maccarone MC, De Trane S, et al. Therapeutic exercise and conservative injection treatment for early knee osteoarthritis in athletes: a scoping review. *Medicina* 2022; 58: 69.

44. Roos EM, Risberg MA and Little CB. Prevention and early treatment, a future focus for OA research. *Osteoart Cart* 2021; 29: 1627–1629.

45. Moons KG, Kengne AP, Grobbee DE, et al. Risk prediction models: II – external validation, model updating, and impact assessment. *Heart* 2012; 98: 691–698.

46. Altman DG, Vergouwe Y, Royston P, et al. Prognosis and prognostic research: validating a prognostic model. *BMJ* 2009; 338: b605.