



Computer-assisted timber identification based on features extracted from microscopic wood sections

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ABSTRACT

Wood anatomy is one of the most important methods for timber identification. However, training wood anatomy experts is time-consuming, while at the same time the number of senior wood anatomists with broad taxonomic expertise is declining. Therefore, we want to explore how a more automated, computer-assisted approach can support accurate wood identification based on microscopic wood anatomy. For our exploratory research, we used an available image dataset that has been applied in several computer vision studies, consisting of 112 — mainly neotropical — tree species representing 20 images of transverse sections for each species. Our study aims to review existing computer vision methods and compare the success of species identification based on (1) several image classifiers based on manually adjusted texture features, and (2) a state-of-the-art approach for image classification based on deep learning, more specifically Convolutional Neural Networks (CNNs). In support of previous studies, a considerable increase of the correct identification is accomplished using deep learning, leading to an accuracy rate up to 95.6%. This remarkably high success rate highlights the fundamental potential of wood anatomy in species identification and motivates us to expand the existing database to an extensive, worldwide reference database with transverse and tangential microscopic images from the most traded timber species and their look-a-likes. This global reference database could serve as a valuable future tool for stakeholders involved in combatting illegal logging and would boost the societal value of wood anatomy along with its collections and experts.

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INTRODUCTION

Progress in computer and internet technology is advancing next-generation phenomics, with innovations in computer science augmenting expert-based description, analysis, and comparison of phenotypes (Houle *et al.* 2010; MacLeod *et al.* 2010; Jordan & Mitchell 2015; Havlíček *et al.* 2019). During the last couple of years, computer vision technologies have boosted the development of tools for the automated description and identification of a wide array of biological objects, including amongst others the shape of leaves (Kumar *et al.* 2012; Wilf *et al.* 2016; Barré *et al.* 2017), herbarium specimens (Unger *et al.* 2016), wings of insects (Favret & Sieracki 2016), flying birds (Antanbori *et al.* 2016), shark fins (Hughes & Burghardt 2016) and animals caught in camera-traps (Gomez Villa *et al.* 2017). In this paper, we focus on a computer-assisted approach for the identification of wood samples from trees and suggest that this identification tool could help protect forests in the future.

Forests cover 30% of the land area on Earth, representing ca. four billion hectares and three trillion trees (Crowther et al. 2015; FAO 2015). The impact on forest net loss during the last 15 years — comparable to the area of France, Spain and the UK combined — has aroused great concern due to the potential loss of a range of ecosystem services, amongst others carbon storage and associated climatic feedbacks, conservation of biodiversity, public recreation, and medicinal products (Bonan 2008; FAO 2015). A detailed assessment of global forest change based on high-resolution satellite imaging shows that deforestation is occurring at an even more disturbing speed, especially in the tropics (Hansen *et al.* 2013; Finer *et al.* 2018). Unsustainable agriculture, mining, and illegal logging contribute to these alarming deforestation rates, representing a massive threat to global biodiversity. It is estimated that more than 100 million m³ of timber are harvested illegally each year, worth between US \$30 and \$100 billion per year (Nelleman 2012; UNODC Committee 2016), but prosecution of illegal logging crimes is hampered by the limited availability of forensic timber identification tools (Dormontt et al. 2015). This is especially true for DNA-based methods, because wood mainly consists of dead fibres and hollow conduits that have lost their living cell contents during cell maturation, and various wood-based treatments like heating, drying and ageing break down the DNA content of the remaining living wood cells (Jiao et al. 2012, 2015). Nevertheless, it remains possible to extract DNA from the sapwood of trees to sequence DNA barcodes for species identification (Jiao et al. 2014, 2018; Nithaniyal et al. 2014), assess geographic provenance (Jolivet & Degen 2012; Vlam et al. 2018), and to reconstruct DNA fingerprints verifying the intact chain of custody for routine trade (Lowe et al. 2010).

There is a considerable body of literature on non-genetic methods which have been successfully applied to identify wood samples: wood anatomy (InsideWood 2004–onwards; Hermanson & Wiedenhoeft 2011; Koch *et al.* 2011; Sarmiento *et al.* 2011; Helmling *et al.* 2018), chemical profiling based on mass spectrometry (Espinoza *et al.* 2015; McClure *et al.* 2015; Evans *et al.* 2017), near-infrared spectroscopy (Russ *et al.* 2009; Braga *et al.* 2011; Pastore *et al.* 2011; Comparison of the spectroscopy (Russ *et al.* 2009; Braga *et al.* 2011; Pastore *et al.* 2011; Pastore *et al.* 2011; Pastore *et al.* 2011; Comparison of the spectroscopy (Russ *et al.* 2009; Braga *et al.* 2011; Pastore *et al.* 2011

al. 2011; Bergo et al. 2016), and detector dogs (Braun 2013). Of these non-genetic tools, wood anatomy remains the most frequently used method for taxonomic identification, as highlighted in the recent UN report on the Best Practice Guide on Forensic Timber Identification (UNODC Committee 2016). However, years of wood anatomical training are required to become an expert, and the number of senior wood anatomists with vast taxonomic knowledge is declining. Likewise, wood anatomy generally allows identification of woods at the genus — not species — level (Gasson 2011), while CITES regulations often require species identification. For instance, only the Malagasy species of the ebony wood genus Diospyros are CITES protected, but wood anatomists are not able to distinguish the protected from the non-protected ebony woods. In an attempt to solve these problems, wood anatomists and computer scientists have been joining forces during the last decade, to come up with ways to strengthen the power of identification through wood anatomy to combat illegal logging (Hermanson & Wiedenhoeft 2011). Some computer-assisted identification tools based on wood anatomy — such as InsideWood, macroHolzdata, CITESwoodID, Pl@ntWood and a new softwoods identification system — are already available (InsideWood 2004onwards; Hermanson & Wiedenhoeft 2011; Koch et al. 2011; Sarmiento et al. 2011; Richter & Dallwitz 2016), and the first portable applications based on macroscopic imaging have been developed to assist customs officers in the field, such as xylotron (https://www.fs. fed.us/research/highlights/highlights display.php?in high id=585), xvlorix (https://www. xylorix.com/), and MyWood-ID (http://mywoodid.frim.gov.my/). However, more sophisticated systems, based on morphometric analyses of cell shapes and mathematical analysis of texture patterns based on an extensive worldwide wood image reference dataset must be further developed in order to allow global species identification in ways unavailable so far.

There have been several studies focusing on computer-assisted wood identification based on transverse macroscopic wood images from a broad taxonomic set of timber species using an array of classical computer vision algorithms. For instance, Khalid *et al.* (2008) showed a high rate of image accuracy (> 95%) based on 20 different Malaysian forest species, and Paula Filho *et al.* (2014) found a recognition rate of up to 88% based on a single texture descriptor using 41 neotropical timber species. A more recent study applied a Convolutional Neural Network approach to macroscopic wood images from 10 species belonging to six genera of the Meliaceae family, with an image recognition success rate from 87% (at species level) to 97% (at genus level; Ravindran *et al.* 2018).

For our research, we have taken advantage of the most extensive, freely available dataset of light microscopic images of transverse wood sections with a sufficient number of replicates per species. This dataset consists of 112 — mainly neotropical — tree species in which each species is represented by 20 microscopic images from transverse sections (Martins *et al.* 2013). We prefer microscopic over macroscopic images because the former includes much more detail that is required to fully explore the value of wood anatomy as an identification tool. We compare different image classifiers based on manually adjusted texture features with a more advanced method based on Convolutional Neural Networks (CNNs) to evaluate computer-assisted microscopic wood anatomy for species identification, and to discuss its potential in the fight against illegal logging. Our results are compared with

results from previous computer vision studies that used the same database to classify all the 2240 images in the Martins database.

MATERIALS AND METHODS

Martins et al. (2013) database

We have used the Martins *et al.* (2013) database, based on pictures from wood anatomical slides generated in the Laboratory of Wood Anatomy at the Federal University of Parana (UFPR) in Curitiba, Brazil. The wood sections were made following the standardized sliding microtome technique, coloured with a solution of acridine red, chrysoidine and astra blue, mounted between glass slides, and subsequently photographed with an Olympus Cx40 microscope equipped with a digital camera generating pictures with a resolution of 1024 \times 768 pixels. The database includes 20 transverse microscopic pictures from 37 gymnosperm species (softwoods) and 75 angiosperm species (hardwoods), in total amounting to 2240 pictures for 112 species. The 20 images per species come from different individuals, but whether or not each sample is derived from another individual, covering the entire distribution range of the species, could not be traced (Luiz Eduardo S. Oliveira, personal communication). Therefore, it is possible that the intraspecific variation captured by the 20 images per species is underestimated for some of the species. The taxonomically cleaned species list is shown in Appendix A, with reference to the phylogenetic position, and its inclusion in CITES and the top 100 most traded timbers (UNODC 2016, Annex 11).

Since the original images of the Martins database are not very high quality, we decided to include a small side project to provisionally assess the impact of better-quality images on the performance of the texture features. Therefore, we made 162 original, higher quality images from transverse sections of 20 tree species in the Martins dataset for which a minimum number of slides per species were available in the wood slide collection of Naturalis Biodiversity Center (see species marked with an asterisk in Appendix A). Most Naturalis images were acquired with a 10× lens (NA 0.75). This results in 24-bit colour images of 2592×1944 pixels representing a field of view of 1.3×1 mm of wood tissue. These Naturalis images were only used to test the classifiers that we produced. The number of species and images in this additional Naturalis test set was used to investigate how well these higher quality images would perform in the classifier, while the original dataset was used to build the classifiers.

Wood identification based on five classifiers and four sets of features

In the jargon of machine learning and especially computational classifiers, we refer to tools based on supervised statistical methods that can be used to segregate a set of images into two or more categories, i.e. species. These species are represented by images of their wood anatomy, and a classifier categorises in such a way that features of different images within each species have similar values, while features of images belonging to different species have contrasting values. To enable classifying the images into the different species, a classifier needs to be trained. This is done by developing a function that optimises the segregation of these images based on features that are extracted from these images. Once the classifier is trained, it presents high precision — meaning accurate recognition of species — when new images are submitted to the trained classifier. This works under the assump-

tion that these new images belong to the same universe of images compared to the images used in the training dataset. It is therefore essential for a good species recognition classifier that the training dataset embodies the natural intraspecific variation of mature wood, which is covered here by 20 images per species.

To develop an idea about the performance of species classification based on the Martins *et al.* (2013) wood dataset, we tested five common classifier strategies together with four manually selected types of features (Table 1). The five classifiers employ different strategies to build the segregating function based on the extracted features: (1) Support Vector Machines identify the function that maximises the distance between dissimilar images; (2) Nearest Neighbour classifiers identify the function that minimises the distance between similar images; (3) Random Forests are based on the construction of multiple classifiers (decision trees aka the forest) using different subsets of the available features

Feature/strategy	Classifier	% recognition
LBP	SVM	89.3
GLCM	SVM	21.4
HOG	SVM	19.3
Gabor	SVM	54.8
LBP	kNN	76.3
GLCM	kNN	35.9
HOG	kNN	8.7
Gabor	kNN	47.2
LBP	RF	76.5
GLCM	RF	40.4
HOG	RF	27.8
Gabor	RF	67.6
LBP	MLP	84.9
GLCM	MLP	37.7
HOG	MLP	13.3
Gabor	MLP	15.1
LBP	LR	89.1
GLCM	LR	29.1
HOG	LR	13.1
Gabor	LR	53.9

Systematic test of the performance of four different texture features for five different classifiers on all 112 species.

In the table, the accuracy per feature set/classifier is given. The Local Binary Pattern (LBP) feature with the Support Vector Machine classifier (first row) clearly presents the best score. Gabor, Gabor feature; GLCM, Grey-level Co-occurrence Matrix; HOG, Histogram of Oriented Gradient feature; LBP, Local Binary Patterns; SVM, Support Vector Machine (linear); kNN, k-Nearest Neighbour; RF, Random Forest; MLP, Multi-Layer Perceptron; LR, Logistic Regression. For all cases tested, 112 classes (species) were in the training and the test set.

Table 1.

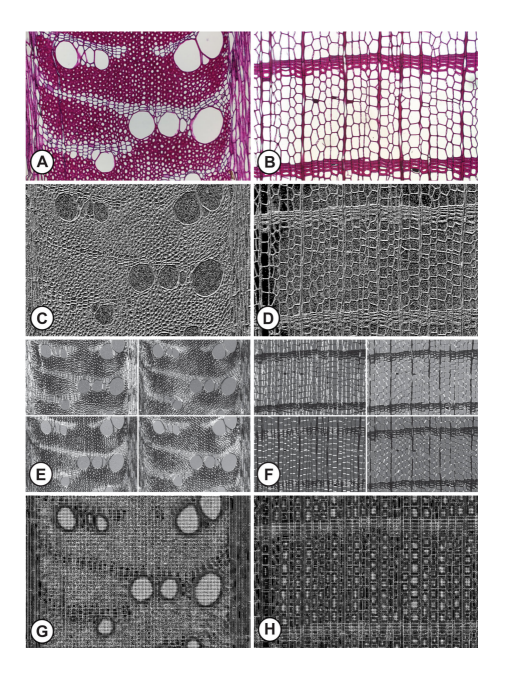
extracted from images; the obtained classifiers are then applied to a new image, and a final decision about the classification of the new image is built based on the aggregation of the results of the different classifiers — for example, by selecting the classification that is produced equally by the largest number of classifiers; (4) Multi-layer Perceptrons employ a stacked collection of linear segregators, aka perceptrons, in order to simulate possibly non-linear segregators; and (5) Logistic Regression classifiers identify the likelihood of an image belonging to a category based on sigmoid non-linear functions. For each classifier, we extracted four features. These features correspond to different image properties: (1) Local Binary Patterns (LBP) correspond to colour and grey-level fluctuations found in a predefined window, i.e. neighbourhood, within a given image (Fig. 1C–D); (2) Grey-Level Co-occurrence Matrices (GLCM) correspond to grev-level fluctuations found in an image as a whole, thus characterising textures; and (3) Gabor features (Fig. 1E–F) and (4) Histogram of Oriented Gradients (HOG; Fig. 1G-H) characterise the gradient of variation in grey-level and colour in image pixels belonging to delimited regions within an image (pictures of Fig. 1 and additional images can also be consulted via http://bio-imaging.liacs.nl/ galleries/IAWA-WoodClassification/).

The aim of a classifier is to predict the species identity of an "unseen" sample. Therefore, the classifier needs to be validated in order to prevent overfitting or selection bias. We have made randomly generated training sets with 10-fold cross-validation for all classifiers. In other words, the image dataset was randomly split into 10 subsets, and for each classifier, the following procedure was used to optimise its performance: (1) one subset was selected, (2) the classifier was trained to segregate the target classes (i.e. wood images) using images in the remaining 9 subsets, (3) the performance of the optimised classifier was tested on the selected subset, (4) steps (1)–(3) were repeated for all of the 10 subsets, in order to obtain an average performance of the classifier was incrementally adjusted to reach optimal average performance.

Average values for accuracy from this cross-validation are reported. The accuracy rates resulting from our classifiers (Table 1) can be compared with the rates in Table 2 that summarises results of previously published papers based on the same Martins dataset, enabling us to evaluate the accuracy of comparable classifiers applying manually adjusted texture features. Details of the computational classifiers used, and their extracted features are available upon request.

Wood identification based on Convolutional Neural Networks

Recent developments in pattern recognition have brought about new evolution to neural networks in the form of Convolutional Neural Networks (CNNs). CNNs advance on multi-layer perceptrons by employing (1) a large collection of layers of perceptrons, such that in each layer the connections between perceptrons are different, and (2) information is transferred between layers based on the mathematical notion of convolution, which combines the output of selected layers in order to build the input for a following layer. With this organisation of perceptrons based on structured connections within layers and convolutions to transfer information across layers, it is possible to combine supervised and un-



Number of	f classes	Features/strategy	Classifier	Recognition	References
Training	Testing			rates (%)	
112	112	LBP	SVM	80.7	Martins et al. (2012)
112	112	LBP + LPQ	SVM	86.5	Martins et al. (2012)
68	44	Ensemble Texture Features	DSC	93	Martins et al. (2015)
112	112	LPQ + GLCM	SVM	93.2	Cavalin <i>et al.</i> (2013)
112	112	LPQ-Blackman	SVM	95	Kapp <i>et al.</i> (2012)
112	112	CNN	CNN	95	Hafemann et al. (2014)

Summary of the results based on computer vision publications that used the Martins *et al.* (2013) dataset of 112 — mainly neotropical — timber species.

Table 2.

Only the last row refers to a deep learning study, the other studies only have used texture-based features. CNN, Convolutional Neural Networks; DSC, Dynamic Selection of Classifier; GLCM, Grey-level Cooccurrence Matrix; LBP, Local Binary Patterns; LPQ, Local Phase Quantization; SVM, Support Vector Machine (Gaussian Kernel).

supervised classification by aggregating images in classes, i.e. unsupervised classification, and then segregating classes according to observed features, i.e. supervised classification. Thus, a CNN can consist of different layers and each of these layers can be connected in different ways. The layers and the connections are the mark-up of the "net" and this structure is referred to as the architecture. So, different architectures can be used to organise perceptrons in CNNs, and each of these architectures has specific characteristics that work well only for specific datasets and require the data to be prepared in specific ways.

The preparation of the data before classification is an important step in CNN. The notion of transfer learning (Pan & Yang 2010), which corresponds to bootstrapping the training steps using images from a different domain — i.e. not related to wood anatomy but featuring similar structure when compared with the anatomical images — has greatly enhanced the learning outcome of the training of classifiers with CNN in our experiments. Using this technique, we have pre-trained the CNN in our experiments with a much larger dataset that is available in *ImageNet* (Russakovsky *et al.* 2013), a publicly available resource of labelled images which is often used for benchmarking of algorithms (http://www.image-net.org/). This helps to avoid training the CNN from scratch with a dataset that is relatively small. In addition, there are strategies in CNN to prevent overfitting by tuning the weights in the

Figure 1. (A) Cross-section of *Grevillea robusta* (angiosperm) and (B) *Taxodium distichum* (gymnosperm), two species included in the Martins database, with visualization of some of the computer extracted features used to classify the timbers. All images are generated using the MatLab Image Processing Toolbox. (A–B) Original cross-sections. (C–D) The Local Binary Pattern (LBP) texture feature maps from A–B; the window size for the LBP computation was 3×3 pixels. (E–F) Gabor file with 4 orientations and a scale of 4; the 4 orientations are assembled in one 4×4 panel (0, 45, 90, 135 orientation, respectively). (G–H) The Histogram of Oriented Gradient feature for A–B; a window size of 8×8 pixels was used to assemble the orientation of the gradients.

Table 3.

Systematic test of the performance of Convolutional Neural Networks based on four different architectures (GoogLeNet, Alexnet, VGG16, ResNet101).

	GoogLeNet	Alexnet	VGG16	ResNet101
Gymno vs angio	96.3	99.2	99.8	99•7
species	80.1	85.9	90.7	96.4

First, a two-class problem (gymnosperms vs angiosperms) was evaluated. Second, all images were evaluated at the species level (112 species). In terms of accuracy, we observed that ResNet101 is the best performer, while GoogLeNet is the worst.

layers of the CNN. These strategies structurally enhance the learning outcome of the CNN, and the performance can still increase further when adding more labelled data. Details of the deep learning analyses are provided in Appendix B.

The results obtained with CNN have been outstanding; however, the deployment of a CNN is not out-of-the-box. We have investigated four popular CNN architectures, namely GoogLeNet, AlexNet, VGG16 and ResNet101, and explored their accuracy with respect to the Martins database. For Alexnet the images were resampled (256×256 pixels), but original images were used for the other architectures. The results of our experiments for the four CNN architectures are listed in Table 3.

RESULTS AND DISCUSSION

We have evaluated the performance of computer vision classification for tree species at the species level based on the largest available database of microscopic images from transverse sections, covering 20 images for 112 timber species (Martins *et al.* 2013). Using a combination of five different classifiers extracting four features each, we conclude that the Support Vector Machine (SVM), using the Local Binary Pattern feature (LBP; Fig. 1C–D), has the best performance to identify the wood images, leading to 89.3% accuracy (Table 1). This surprisingly high recognition rate based on only transverse sections outperforms the LBP analysis in the initial Martins *et al.* (2012) experiments (89.3 vs 80.7%; Tables 1 and 2). This is due to the experimentation with the parameters of the LBP method: given the resolution of the image, there is an optimum that can be accomplished for the feature to come out best.

In the past, the Martins dataset has been subjected to the extraction of combinations of additional features to further increase the species recognition rate (Table 2). For instance, the combination of Local Binary Patterns and Local Phase Quantization features initially yielded a recognition rate of 86.5% (Martins *et al.* 2012). Martins *et al.* (2015) applied a large collection of texture-based features to explore local patterns found in images, namely Gabor filters and Local Phase Quantization, and point-based features to explore individual pixel values — namely scale-invariant feature transform (SIFT) and speed-up robust feature (SURF) — resulting in a similar recognition score of 93%. Another approach, referred to as the quadtree approach resulting in a regular subdivision in square regions within the image (Samet 1984), was applied by Cavalin *et al.* (2013). Computing an ensemble of features in these subdivisions resulted in a boosting of the recognition rate to 93.2%. Kapp *et*

al. (2012) also used a quadtree-based approach to assess multiple sets of features on the same dataset and reported recognition scores of 95%. Finally, Hafemann *et al.* (2014) compared textural descriptors with classical classifiers to the performance of a Convolutional Neural Network (CNN), which yielded an accuracy of 95% (Table 2).

Based on CNN with ResNet101 architecture in our study, we could further improve this high species recognition rate in our study up to 96.4%, while at the gymnosperm versus angiosperm level a recognition rate of over 99% could be accomplished for all three CNN architectures in our evaluation (Table 3). Misidentification at the species level may have been caused by the striking resemblance between transverse sections of the same species, poor quality of some of the sections, limits to image quality in the dataset, or by taxonomic synonymy issues. For example, the original Martins dataset included *Cephalotaxus dru*pacea and C. harringtonia as separate species, while these are now considered as synonyms (Appendix A). This implies that the species recognition rates in Tables 1–3 should be even higher because some of the mismatches (false negatives) are actually classified correctly (true positives). The few misidentifications at the higher gymnosperm vs angiosperm level is due to the inclusion of *Ephedra*, which is one of the few vessel-bearing gymnosperms that resemble more the vessel-bearing angiosperms compared to the typical tracheid-based gymnosperm wood in transverse sections. Further insight in classification performance requires the images to be of excellent quality, which is not the case for the original images in the Martins database. We have learned from our preliminary analysis based on the original, higher resolution Naturalis images that the texture features studied resulted in higher performance scores over the complete spectrum of features. However, for a classifier to work well with the self-generated images, we could not add a sufficient number of images to support this. This means that we only used the Naturalis images in testing. Further experiments are required to define how many images per species need to be added to get a stable and accurate prediction from the trained classifier, based on the quality of the images. In future work, we will also combine our results with ongoing research about the combination of knowledge-based reasoning and machine learning to improve the performance of the classifiers, enabling us to account explicitly for expert knowledge together with knowledge synthesis obtained through machine learning.

The high species recognition scores presented in Tables 1–3 support the validity of wood anatomy in species identification and encourage us to further expand the existing Martins dataset with extra microscopic images. As a first step, it would be interesting to evaluate how the species recognition rates would be further improved when a complementary dataset of tangential wood images (20 per species, preferably from 20 mature individuals covering the entire range of the species) is added. As a second step, building a more complete reference database, including also transverse and tangential images from the top 100 traded timber species (annex 11 in UNODC Committee 2016) and all the CITES-listed timbers and their look-a-likes, is desired to further finetune existing computer-assisted tools to deal with more challenging classifying problems. Adding radial sections is not desired as the length of the rays in radial sections heavily relies on the exact radial orientation plane and the waviness of the rays, thereby jeopardizing the accuracy of the classifier.

In terms of the workflow for the extension of the dataset, it is of vital importance that a wood anatomist makes a first assessment on the representativeness of a wood sample or an available glass slide. The online information provided by scientific wood collections around the world summarised in Index Xylariorum (Stern 1988) and continuously updated under the guidance of IAWA (http://www.iawa-website.org/en/Wood collection/Overview of Wood Collection.shtml) and GTTN (https://globaltimbertrackingnetwork.org/products/ iawa-index-xylariorum/), is helpful in this regard. Once collected and approved, the wood sample can be further processed (sections and images), and subsequently added to the dataset. In order to understand texture features, the resolving power of the image should be properly addressed and taken into account at the acquisition. For example, the underperformance of some of the features (Tables 1 and 2) can be partially understood by the lack of resolution — and thus detail — of some of the images. Therefore, the process of selecting wood samples and making high-quality images should be carried out in close collaboration between a wood anatomist and a computational scientist with thorough knowledge in imaging and machine learning. In this manner, we can produce a high-quality dataset for the training of the classifier.

If these global wood databases would be matched with complementary databases (anatomy, chemical profiling, DNA) representing the same species, wood scientists can develop a robust tool that facilitates (1) identification of traded woods in an efficient and accurate way, (2) enables fine-scale assessment of geographic provenance of imported logs, and (3) allows DNA fingerprinting. These complementary tools would boost forensic timber research and permit courts to prosecute wood trading companies for smuggling illegal timber. Examples of detrimental fines have already been imposed by the court to wood trading companies (e.g., Lumber Liquidators, Gibson Guitars), proving that implementation of timber laws (e.g., EUTR, Lacey act) can be successful. To make the anatomy tool generally accessible, the trained computer algorithms can be implemented in a cloud-based, computer vision tool enabling everyone to identify illegally traded timbers. However, this can only work when front-line officers and other stakeholders in the field know how to prepare (unstained) wood sections with sufficient quality and upload pictures to the wood identification web server. Also, here wood anatomists can play an important role in training stakeholders using proper basic tools (cooking pot, sliding microtome, sharp disposable blades, standard light microscope with camera). Evidently, before deciding to make wood sections, stakeholders can do a preliminary screening using one of the available portable devices to identify woods based on a cut surface (e.g. xylotron, xylorix), and then continue making wood sections of the suspicious samples.

CONCLUSIONS

The use of traditional wood anatomical descriptions following standardized features (IAWA Committee 1989, 2004) has been successfully applied to characterize and compare angiosperm and gymnosperm genera, but we are facing its limits with respect to timber identification at the species level. Computer-assisted classification based on texture features takes into account the standardized anatomical features and at the same time integrates other features like colour variation and acuteness of edges. We, therefore, believe

that the use of computer vision techniques presents a powerful tool to boost the accuracy of timber recognition based on microscopic wood anatomy. In this paper, we found a remarkably high success rate in computer-assisted identification using only transverse wood sections from 112 species (Tables 1–3). However, we must stress that this Martins database is far from complete as a test database to rigorously assess the value of computer vision algorithms in timber identification. Therefore, we encourage expanding this database by adding wood samples from the top 100 most traded timber species, along with the CITESlisted timber species and their look-a-likes. Using close interaction between computational scientists and wood anatomists, our community can compile an extensive, worldwide reference database including a sufficient number of transverse and tangential microscopic images from the species of interest. Whether or not 20 images per species will suffice, is a topic for further investigation and will probably also depend on various image-related aspects, such as image resolution, biological and staining variation amongst the images, and section artefacts.

In the short term, it will be important to develop this global reference database of microscopic wood images representing all relevant traded timber species. In addition, training of classifiers should be benchmarked on a large image dataset in combination with analyses of the mismatches, as only then the true benefits of the classification procedure emerge. In the longer term, a global concerted effort between wood anatomists and computer scientists should be able to implement an open-access, cloud-based, computer-vision-based classification tool enabling a more efficient and more accurate identification process of wood. This wood identification tool would benefit scientists, customs officers, and other stakeholders who are confronted with identification of wood samples, including bona fide wood trading companies that are eager to receive more support in efficient identification protocols for their shipments. Moreover, we envision that the accuracy of this online tool will be robust enough to support prosecutions of wood trading companies in court that violate (inter)national regulations, and evidently these online identifications must be validated in a court of law by wood anatomy experts. It would be a great opportunity for IAWA to play a leading role in facilitating both the short- and longer-term goals, and to ideally also act as a coordinator to help building complementary reference datasets based on DNA based methods and chemical profiling. Only a collaborative effort amongst wood biologists will be able to contribute to the conservation of our forests and its wildlife and will generate an increasing awareness of the societal relevance of scientific wood collections and wood anatomical expertise worldwide.

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Overview of the 112 woody species included in the Martins database, with reference to the phylogenetic position, and its inclusion in CITES and the top 100 Species synonymy has updated using the Plants of the World Online portal (POWO 2019; http://www.plantsoftheworldonline.org/). Additional images for transmost traded timbers sensu UNODC (2016).

verse wood sections from available microscopic slides from the Naturalis collection were added to the original Martins database for the 20 species marked with an asterisk (with number of Naturalis pictures in parentheses).

Class	Order	Family	Genus	Species name	CITES protected	Top 100 traded timbers
Gymnosperms	Ginkgoales Pinales	Ginkgoaceae Araucariaceae	Ginkgo Agathis	biloba (10)* Beccarii		
	Pinales	Araucariaceae	Araucaria	Angustifolia		х
	Pinales	Cupressaceae	Calocedrus	Decurrens		
	Pinales	Cupressaceae	Chamaecyparis	Formosensis		
	Pinales	Cupressaceae	Chamaecyparis	$pisifera~(6)^*$		
	Pinales	Cupressaceae	Fitzroya	cupressoides	Appendix I	
	Pinales	Cupressaceae	Hesperocyparis	Arizonica		
	Pinales	Cupressaceae	Hesperocyparis	Lindleyi		
	Pinales	Cupressaceae	Larix	kaempferi		
	Pinales	Cupressaceae	Larix	Laricina		
	Pinales	Cupressaceae	Larix	ds		
	Pinales	Cupressaceae	Sequoia	sempervirens $(15)^*$		
	Pinales	Cupressaceae	Taxodium	$distichum (9)^*$		
	Pinales	Cupressaceae	Tetraclinis	articulata		
	Pinales	Cupressaceae	Widdringtonia	Nodiflora		
	Pinales	Pinaceae	Abies	Religiosa		
	Pinales	Pinaceae	Abies	Vejarii		
	Pinales	Pinaceae	Cedrus	Atlantica		Cedar is listed (unknown sp.)
	Pinales	Pinaceae	Cedrus	libani (5)*		Cedar is listed (unknown sp.)
	Pinales	Pinaceae	Cedrus	ds		Cedar is listed (unknown sp.)

Class	Order	Family	Genus	Species name	CITES protected	Top 1 00 traded timbers
	Pinales	Pinaceae	Keteleeria	Fortune		
	Pinales	Pinaceae	Picea	abies (9)*		х
	Pinales	Pinaceae	Pinus	Arizonica		
	Pinales	Pinaceae	Pinus	Caribaea		
	Pinales	Pinaceae	Pinus	Elliottii		
	Pinales	Pinaceae	Pinus	gregii		
	Pinales	Pinaceae	Pinus	maximinoi		
	Pinales	Pinaceae	Pinus	Taeda		
	Pinales	Pinaceae	Pseudotsuga	Menziesii		
	Pinales	Pinaceae	Tsuga	canadensis		
	Pinales	Pinaceae	Tsuga	sp		
	Pinales	Podocarpaceae	Podocarpus	Lambertii		
	Pinales	Taxaceae	Cephalotaxus	harringtonia (formerly drupacea)		
	Pinales	Taxaceae	Cephalotaxus	harringtonia		
	Pinales	Taxaceae	Taxus	Baccata		х
	Pinales	Taxaceae	Torreya	Nucifera		
	Gnetales	Ephedraceae	Ephedra	californica		
Angiosperms	Magnoliales	Annonaceae	Porcelia	macrocarpa		
	Magnoliales	Magnoliaceae	Magnolia	$grandiflora~(7)^*$		
	Magnoliales	Magnoliaceae	Magnolia	Ovata		
	Magnoliales	Myristicaceae	Bicuiba	Oleifera		
	Laurales	Lauraceae	Nectandra	oppositifolia		
	Laurales	Lauraceae	Nectandra	sp		
	Laurales	Lauraceae	Ocotea	porosa		
	Laurales	Lauraceae	Persea	Venosa		

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APPENDIX A. (Continued.)

Class	Order	Family	Genus	Species name	CITES protected	Top 100 traded timbers
	Proteales	Proteaceae	Grevillea	$robusta~(6)^*$		
	Proteales	Proteaceae	Grevillea	sp		
	Proteales	Proteaceae	Roupala	ds		
	Fabales	Fabaceae	Abarema	jupunba		
	Fabales	Fabaceae	Anadenanthera	colubrina		
	Fabales	Fabaceae	Anadenanthera	peregrina (6)*		
	Fabales	Fabaceae	Copaifera	trapezifolia (6)*		
	Fabales	Fabaceae	Dalbergia	nigra	Appendix I	х
	Fabales	Fabaceae	Dalbergia	frutescens	Appendix II	х
	Fabales	Fabaceae	Dalbergia	spruceana	Appendix II	х
	Fabales	Fabaceae	Dinizia	excelsa		
	Fabales	Fabaceae	Enterolobium	$schomburgkii~(7)^*$		
	Fabales	Fabaceae	Eperua	falcata (11)*		
	Fabales	Fabaceae	Hymenaea	courbaril (9)*		х
	Fabales	Fabaceae	Hymenaea	sp		х
	Fabales	Fabaceae	Inga	sessilis		
	Fabales	Fabaceae	Leucaena	leucocephala		
	Fabales	Fabaceae	Lonchocarpus	subglaucescens		
	Fabales	Fabaceae	Mimosa	bimucronata		
	Fabales	Fabaceae	Mimosa	scabrella		
	Fabales	Fabaceae	Ormosia	excelsa		
	Fabales	Fabaceae	Parapiptadenia	excelsa		
	Fabales	Fabaceae	Parapiptadenia	rigida (6)*		
	Fabales	Fabaceae	Parkia	Multijuga		
	Fabales	Fabaceae	Pterocarpus	Violaceus		

(Continued.)						
Class	Order	Family	Genus	Species name	CITES protected	Top 1 oo traded timbers
	Fabales	Fabaceae	Senegalia	tucunamensis		
	Fabales	Fabaceae	Schizolobium	Parahyba		
	Rosales	Moraceae	Bagassa	guianensis (7)*		
	Rosales	Moraceae	Brosimum	alicastrum $(7)^*$		
	Rosales	Moraceae	Ficus	gomelleira		
	Rosales	Rhamnaceae	Frangula	alnus		
	Rosales	Rhamnaceae	Hovenia	dulcis		
	Rosales	Rosaceae	Prunus	reflexa		х
	Rosales	Rosaceae	Prunus	serotina		х
	Myrtales	Melastomataceae	Tibouchina	sellowiana		
	Myrtales	Myrtaceae	Campomanesia	xantho carpa		
	Myrtales	Myrtaceae	Eucalyptus	globulus (9)*		
	Myrtales	Myrtaceae	Eucalyptus	grandis		х
	Myrtales	Myrtaceae	Eucalyptus	saligna		х
	Myrtales	Myrtaceae	Myrcia	racemulosa		
	Myrtales	Vochysiaceae	Erisma	uncinatum		
	Myrtales	Vochysiaceae	Qualea	ds		
	Myrtales	Vochysiaceae	Vochysia	laurifolia		
	Sapindales	Meliaceae	Cabralea	canjerana		
	Sapindales	Meliaceae	Carapa	guianensis		
	Sapindales	Meliaceae	Cedrela	fissilis	Appendix III	х
	Sapindales	Meliaceae	Khaya	ivorensis $(8)^*$		х
	Sapindales	Meliaceae	Melia	azedarach		
	Sapindales	Meliaceae	Swietenia	macrophylla (11)*	Appendix II	х

Class	Order	Family	Genus	Species name	CITES protected	Top 1 oo traded timbers
	Sapindales	Rutaceae	Balfourodendron	riedelianum		
	Sapindales	Rutaceae	Citrus	xaurantium		
	Sapindales	Rutaceae	Zan tho xylum	rhoifolium		
	Sapindales	Simaroubaceae	Simarouba	amara		
	Ericales	Lecythidaceae	Cariniana	estrellensis		
	Ericales	Lecythidaceae	Couratari	sp		
	Ericales	Lecythidaceae		coriacea		
	Ericales	Lecythidaceae		chartacea		
	Ericales	Sapotaceae		lucentifolium		
	Ericales	Sapotaceae	Chrysophyllum	sp		
	Ericales	Sapotaceae	Micropholis	guyanensis		
	Gentianales	Rubiaceae	Faramea	occidentalis		
	Gentianales	Rubiaceae	Psychotria	carthagenensis		
	Gentianales	Rubiaceae	Palicourea	jungiana		
	Lamiales	Bignoniaceae	Tabebuia	roseo alba		х
	Lamiales	Bignoniaceae	Tabebuia	sp		х
	Lamiales	Oleaceae	Ligustrum	$lucidum (6)^*$		

APPENDIX A. (Continued.)

APPENDIX B: DETAILS OF THE DEEP LEARNING ANALYSES

The different nets for the deep learning classification were implemented in *Caffe* (https:// caffe.berkeleyvision.org/). In addition, we used numpy, scikit-learn, and OpenCV-python in the experiments. For convergent nets, we have used deploy.prototxt to do inference work. The deep nets, i.e., VGGnets, Residual Nets and GoogLeNet, were all run with the same training parameters. Both the Martins and the Naturalis-extended data set were used in the training. The nets were trained on an Ubuntu-based Linux workstation that is equipped with two TITAN GPU NVidia cards.

Within the *Caffe* environment the following settings are used:

>net: "Users/"	# Path to the training network architecture file, specifically train_val.prototxt.
>test_iter: 50	<pre># Number of test/validate iteration. To be noticed, test_size == test_iter *</pre>
	test_batch_size.
test_interval: 500	# Every 500 training iterations, forward one test validation.
base_lr: 0.001	# The original learning rate.
lr_policy: "step"	# The adjustment strategy of learning rate. If set as "step", then base_lr *
	gamma (floor(iter / stepsize)), iter here refers to the current iteration
	number.
gamma: 0.1	# Hyperparameter in the learning rate updating.
stepsize: 1000	# Hyperparameter in the learning rate calculation.
display: 50	# Display training loss and accuracy every 50 iterations.
max_iter: 20000	# The maximum number of training iterations.
momentum: 0.9	# Optimization hyperparameter in weighted gradient descent.
weight_decay: 0.0005	# Hyperparameter in preventing from overfitting.
snapshot: 2000 #	In order to restart the training process quickly, weights and network status
	are saved as a snapshot every 2000 iterations.
snapshot_prefix: " "	# Prefix of saved snapshot models.
solver_mode: GPU	# Using GPUs as computing resources.

Detailed information is available on request. We do realize that the current approach for deep learning has been moved to *Tensorflow* (https://www.tensorflow.org/) and other environments. However, our original experiments were performed with *Caffe*, and therefore we present specifically these parameters.