Trade-off between deep learning for species identification and inference about predator-prey co-occurrence: Reproducible R workflow integrating models in computer vision and ecological statistics

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Abstract

Deep learning is used in computer vision problems with important applications in several scientific fields. In ecology for example, there is a growing interest in deep learning for automatizing repetitive analyses on large amounts of images, such as animal species identification.

However, there are challenging issues toward the wide adoption of deep learning by the community of ecologists. First, there is a programming barrier as most algorithms are written in Python while most ecologists are versed in R. Second, recent applications of deep learning in ecology have focused on computational aspects and simple tasks without addressing the underlying ecological questions or carrying out the statistical data analysis to answer these questions.

Here, we showcase a reproducible R workflow integrating both deep learning and statistical models using predator-prey relationships as a case study. We illustrate deep learning for the identification of animal species on images collected with camera traps, and quantify spatial co-occurrence using multispecies occupancy models.

Despite average model classification performances, ecological inference was similar whether we analysed the ground truth dataset or the classified dataset. This result calls for further work on the trade-offs between time and resources allocated to train models with deep learning and our ability to properly address key ecological questions with biodiversity monitoring. We hope that our reproducible workflow will be useful to ecologists and applied statisticians.

All material (source of the Rmarkdown notebook and auxiliary files) is available from https://github.com/oli_viergimenez/computo-deeplearning-occupany-lynx.

Introduction

Computer vision is a field of artificial intelligence in which a machine is taught how to extract and interpret the content of an image (Krizhevsky, Sutskever, and Hinton 2012). Computer vision relies on deep learning that allows computational models to learn from training data – a set of manually labelled images – and make predictions on new data – a set of unlabelled images (Baraniuk, Donoho, and Gavish 2020; LeCun, Bengio, and Hinton 2015). With the growing availability of massive data, computer vision with deep learning is being increasingly used to perform tasks such as object detection, face recognition, action and activity recognition or human pose estimation in fields as diverse as medicine, robotics, transportation, genomics, sports and agriculture (Voulodimos et al. 2018).

In ecology in particular, there is a growing interest in deep learning for automatizing repetitive analyses on large amounts of images, such as identifying plant and animal species, distinguishing individuals of the same
or different species, counting individuals or detecting relevant features (Christin, Hervet, and Lecomte 2019; Lamba et al. 2019; Weinstein 2018). By saving hours of manual data analyses and tapping into massive amounts of data that keep accumulating with technological advances, deep learning has the potential to become an essential tool for ecologists and applied statisticians.

Despite the promising future of computer vision and deep learning, there are challenging issues toward their wide adoption by the community of ecologists (e.g. Wearn, Freeman, and Jacoby 2019). First, there is a programming barrier as most, if not all, algorithms are written in the Python language while most ecologists are versed in R (Lai et al. 2019). If ecologists are to use computer vision in routine, there is a need for bridges between these two languages (through, e.g., the reticulate package Allaire et al. (2017) or the shiny package Tabak et al. (2020)). Second, recent applications of computer vision via deep learning in ecology have focused on computational aspects and simple tasks without addressing the underlying ecological questions (Sutherland et al. 2013), or carrying out statistical data analysis to answer these questions (Gimenez et al. 2014). Although perfectly understandable given the challenges at hand, we argue that a better integration of the why (ecological questions), the what (automatically labelled images) and the how (statistics) would be beneficial to computer vision for ecology (see also Weinstein 2018).

Here, we showcase a full why-what-how workflow in R using a case study on the structure of an ecological community (a set of co-occurring species) composed of the Eurasian lynx (Lynx lynx) and its two main preys. First, we introduce the case study and motivate the need for deep learning. Second we illustrate deep learning for the identification of animal species in large amounts of images, including model training and validation with a dataset of labelled images, and prediction with a new dataset of unlabelled images. Last, we proceed with the quantification of spatial co-occurrence using statistical models.

### Collecting images with camera traps

Lynx (Lynx lynx) went extinct in France at the end of the 19th century due to habitat degradation, human persecution and decrease in prey availability (Vandel and Stahl 2005). The species was reintroduced in Switzerland in the 1970s (Breitenmoser 1998), then re-colonised France through the Jura mountains in the 1980s (Vandel and Stahl 2005). The species is listed as endangered under the 2017 IUCN Red list and is of conservation concern in France due to habitat fragmentation, poaching and collisions with vehicles. The Jura holds the bulk of the French lynx population.

To better understand its distribution, we need to quantify its interactions with its main preys, roe deer (Capreolus capreolus) and chamois (Rupicapra rupicapra) (Molinari-Jobin et al. 2007), two ungulate species that are also hunted. To assess the relative contribution of predation and hunting, a predator-prey program was set up jointly by the French Office for Biodiversity, the Federations of Hunters from the Jura, Ain and Haute-Savoie counties and the French National Centre for Scientific Research.

Animal detections were made using a set of camera traps in the Jura mountains that were deployed in the Jura and Ain counties (see Figure 1). We divided the two study areas into grids of $2.7 \times 2.7$ km cells or sites hereafter (Zimmermann et al. 2013) in which we set two camera traps per site (Xenon white flash with passive infrared trigger mechanisms, model Capture, Ambush and Attack; Cuddeback), with 18 sites in the Jura study area, and 11 in the Ain study area that were active over the study period (from February 2016 to October 2017 for the Jura county, and from February 2017 to May 2019 for the Ain county). The location of camera traps was chosen to maximise lynx detection. Camera traps were checked weekly to change memory cards, batteries and to remove fresh snow after heavy snowfall.

In total, 45563 and 18044 pictures were considered in the Jura and Ain sites respectively after manually dropping empty pictures and pictures with unidentified species. Note that classifying empty images could be automated with deep learning (Norouzzadeh et al. 2021; Tabak et al. 2020). We identified the species present on all images by hand (see Table 1) using digiKam a free open-source digital photo management application (https://www.digikam.org/). This operation took several weeks of labor full time, which is often identified as a limitation of camera trap studies. To expedite this tedious task, computer vision with deep learning has been identified as a promising approach (Norouzzadeh et al. 2021; Tabak et al. 2019; Willi et al. 2019).
Figure 1: Study area, grid and camera trap locations.

Table 1: Species identified in the Jura and Ain study sites with samples size (n). Only first 10 species with most images are shown.

| Species in Jura study site | n   | Species in Ain study site | n   |
|----------------------------|-----|---------------------------|-----|
| human                      | 31644| human                     | 4946|
| vehicle                    | 5637 | vehicle                   | 4454|
| dog                        | 2779 | dog                       | 2310|
| fox                        | 2088 | fox                       | 1587|
| chamois                    | 919  | rider                     | 1025|
| wild board                 | 522  | roe deer                  | 860 |
| badger                     | 401  | chamois                   | 780 |
| roe deer                   | 368  | hunter                    | 593 |
| cat                        | 343  | wild board                | 514 |
| lynx                       | 302  | badger                    | 461 |
Table 2: Model training performance. Images from the Jura study site were used for training.

| species   | precision | recall |
|-----------|-----------|--------|
| badger    | 0.78      | 0.88   |
| red deer  | 0.67      | 0.21   |
| chamois   | 0.86      | 0.81   |
| cat       | 0.89      | 0.78   |
| roe deer  | 0.67      | 0.81   |
| dog       | 0.78      | 0.84   |
| human     | 0.99      | 0.79   |
| hare      | 0.32      | 0.52   |
| lynx      | 0.87      | 0.95   |
| fox       | 0.85      | 0.90   |
| wild boar | 0.93      | 0.88   |
| vehicle   | 0.95      | 0.98   |

Deep learning for species identification

Using the images we obtained with camera traps (Table 1), we trained a model for identifying species using the Jura study site as a calibration dataset. We then assessed this model’s ability to automatically identify species on a new dataset, also known as transferability, using the Ain study site as an evaluation dataset.

Training - Jura study site

We selected at random 80% of the annotated images for each species in the Jura study site for training, and 20% for testing. We applied various transformations (flipping, brightness and contrast modifications; Shorten and Khoshgoftaar (2019)) to improve training (see Appendix). To reduce model training time and overcome the small number of images, we used transfer learning (Yosinski et al. 2014; Shao, Zhu, and Li 2015) and considered a pre-trained model as a starting point. Specifically, we trained a deep convolutional neural network (ResNet-50) architecture (He et al. 2016) using the fastai library (https://docs.fast.ai/) that implements the PyTorch library (Paszke et al. 2019). Interestingly, the fastai library comes with an R interface (https://eagerai.github.io/fastai/) that uses the reticulate package to communicate with Python, therefore allowing R users to access up-to-date deep learning tools. We trained models on the Montpellier Bioinformatics Biodiversity platform using a GPU machine (Titan Xp nvidia) with 16Go of RAM. We used 20 epochs which took approximately 10 hours. The computational burden prevented us from providing a full reproducible analysis, but we do so with a subsample of the dataset in the Appendix. All trained models are available from https://doi.org/10.5281/zenodo.5164796.

We calculated three metrics to evaluate our model performance at correctly identifying species (e.g. Duggan et al. 2021). Specifically, we relied on accuracy the ratio of correct predictions to the total number of predictions, recall a measure of false negatives (FN; e.g. an image with a lynx for which our model predicts another species) with recall = TP / (TP + FN) where TP is for true positives, and precision a measure of false positives (FP; e.g. an image with any species but a lynx for which our model predicts a lynx) with precision = TP / (TP + FP). In camera trap studies, a strategy (Duggan et al. 2021) consists in optimizing precision if the focus is on rare species (lynx), while recall should be optimized if the focus is on common species (chamois and roe deer).

We achieved 85% accuracy during training. Our model had good performances for the three classes we were interested in, with 87% precision for lynx and 81% recall for both roe deer and chamois (Table 2).

Transferability - Ain study site

We evaluated transferability for our trained model by predicting species on images from the Ain study site which were not used for training. Precision was 77% for lynx, and while we achieved 86% recall for roe deer,
our model performed poorly for chamois with 8% recall (Table 3).

To better understand this pattern, we display the results under the form of a confusion matrix that compares model classifications to manual classifications (Figure 2). There were a lot of false negatives for chamois, meaning that when a chamois was present in an image, it was often classified as another species by our model.

Overall, our model trained on images from the Jura study site did poorly at correctly predicting species on images from the Ain study site. This result does not come as a surprise, as generalizing classification algorithms to new environments is known to be difficult (Beery, Horn, and Perona 2018). While a computer scientist might be disappointed in these results, an ecologist would probably wonder whether ecological inference about the interactions between lynx and its prey is biased by these average performances, a question we address in the next section.

**Spatial co-occurrence**

Here, we analysed the data we acquired from the previous section. For the sake of comparison, we considered two datasets, one made of the images manually labelled for both the Jura and Ain study sites pooled together (*ground truth dataset*), and the other in which we pooled the images that were manually labelled for the Jura study site and the images that were automatically labelled for the Ain study site using our trained model (*classified dataset*).

We formatted the data by generating monthly detection histories, that is a sequence of detections ($Y_{sit} = 1$) and non-detections ($Y_{sit} = 0$), for species $s$ at site $i$ and sampling occasion $t$ (see Figure 3).

To quantify spatial co-occurrence between lynx and its preys, we used a multispecies occupancy modeling approach (Rota et al. 2016; Clipp et al. 2021) using the R package *unmarked* (Fiske and Chandler 2011). The multispecies occupancy model assumes that observations $y_{sit}$, conditional on $Z_{si}$ the latent occupancy state of species $s$ at site $i$ are drawn from Bernoulli random variables $Y_{sit}|Z_{si} \sim \text{Bernoulli}(Z_{si}p_{sit})$ where $p_{sit}$ is the detection probability of species $s$ at site $i$ and sampling occasion $t$. Detection probabilities can be modeled as a function of site and/or sampling covariates, or the presence/absence of other species, but for the sake of illustration, we will make them only species-specific here.

The latent occupancy states are assumed to be distributed as multivariate Bernoulli random variables...
Figure 2: Confusion matrix comparing automatic to manual species classifications. Species that were predicted by our model are in columns, and species that are actually in the images are in rows.
Figure 3: Detections (black) and non-detections (light grey) for each of the 3 species lynx, chamois and roe deer between March and November for all years pooled together. Sites are on the Y axis, while sampling occasions are on the X axis. Only data from the ground truth dataset are displayed.
Let us consider 2 species, species 1 and 2, then \( Z_i = (Z_{i1}, Z_{i2}) \sim \text{multivariate Bernoulli}(\psi_{11}, \psi_{10}, \psi_{01}, \psi_{00}) \) where \( \psi_{11} \) is the probability that a site is occupied by both species 1 and 2, \( \psi_{10} \) the probability that a site is occupied by species 1 but not 2, \( \psi_{01} \) the probability that a site is occupied by species 2 but not 1, and \( \psi_{00} \) the probability a site is occupied by none of them. Note that we considered species-specific only occupancy probabilities but these could be modeled as site-specific covariates. Marginal occupancy probabilities are obtained as \( \Pr(Z_{i1} = 1) = \psi_{11} + \psi_{10} \) and \( \Pr(Z_{i2} = 1) = \psi_{11} + \psi_{01} \). With this model, we may also infer potential interactions by calculating conditional probabilities such as for example the probability of a site being occupied by species 2 conditional of species 1 with \( \Pr(Z_{i2} = 1 | Z_{i1} = 1) = \frac{\psi_{11}}{\psi_{11} + \psi_{10}}. \)

Detection probabilities were indistinguishable whether we used the ground truth or the classified dataset, with \( p_{\text{lynx}} = 0.51(0.45, 0.58), p_{\text{roe deer}} = 0.63(0.57, 0.68) \) and \( p_{\text{chamois}} = 0.61(0.55, 0.67) \).

We also found that occupancy probability estimates were similar whether we used the ground truth or the classified dataset (Figure 4). Roe deer was the prevalent species, but lynx and chamois were also occurring with high probability (Figure 4).

![Figure 4: Marginal occupancy probabilities for all three species, lynx, roe deer and chamois). Parameter estimates are from a multispecies occupancy model using either the ground truth dataset (in red) or the classified dataset (in blue-grey).](image)

Because marginal occupancy probabilities were high, probabilities of co-occurrence were also estimated high (Figure 5). Our results should be interpreted bearing in mind that co-occurrence is a necessary but not sufficient condition for actual interaction. When both preys were present, lynx was more present than when they were both absent (Figure 5). Lynx was more sensitive to the presence of roe deer than that of chamois (Figure 5).
Figure 5: Lynx occupancy probability conditional on the presence or absence of its preys (roe deer and chamois). Parameter estimates are from a multispecies occupancy model using either the ground truth dataset (in red) or the classified dataset (in blue-grey).
Discussion

In this paper, we aimed at illustrating a reproducible workflow for studying the structure of an animal community and species spatial co-occurrence (why) using images acquired from camera traps and automatically labelled with deep learning (what) which we analysed with statistical occupancy models accounting for imperfect species detection (why). Overall, we found that, even though model transferability could be improved, inference about the potential interactions between lynx and its preys was similar whether we analysed the ground truth data or classified data.

This result calls for further work on the trade-offs between time and resources allocated to train models with deep learning and our ability to correctly answer key ecological questions with camera-trap surveys. In other words, while a computer scientist might be keen on spending time training models to achieve top performances, an ecologist would rather rely on a model showing average performances and use this time to proceed with statistical analyses if, of course, errors in computer-annotated images do not make ecological inference flawed. The right balance may be found with collaborative projects in which scientists from artificial intelligence, statistics and ecology agree on a common objective, and identify research questions that can pick the interest of all parties.

Our demonstration remains however empirical, and we encourage others to try and replicate our results. We also see two avenues of research that could benefit the integration of deep learning and ecological statistics. First, a simulation study could be conducted to evaluate bias and precision in ecological parameter estimators with regard to errors in image annotation by computers. The outcome of this exercise could be, for example, guidelines informing on the confidence an investigator may place in ecological inference as a function of the amount of false negatives and false positives. Second, annotation errors could be accommodated directly in statistical models. For example, single-species occupancy models account for false negatives when a species is not detected by the camera at a site where it is present, as well as false positives when a species is detected at a site where it is not present due to species misidentification by the observer (Miller et al. 2011). Pending a careful distinction between ecological vs. computer-generated false negatives and false positives, error rates could be added to multispecies occupancy models (Chambert et al. 2018) and informed by recall and precision metrics obtained during model training (Tabak et al. 2020).

With regard to the case study, our results should be seen only as preliminary. First, we aim at quantifying the relative contribution of biotic (lynx predation on chamois and roe deer) and abiotic (habitat quality) processes to the composition and dynamic of this ecological community. Second, to benefit future camera trap studies of lynx in the Jura mountains, we plan to train a model again using manually annotated images from both the Jura and the Ain study sites. These perspectives are the object of ongoing work.

With the rapid advances in technologies for biodiversity monitoring (Lahoz-Monfort and Magrath 2021), the possibility of analysing large amounts of images makes deep learning appealing to ecologists. We hope that our proposal of a reproducible R workflow for deep learning and statistical ecology will encourage further studies in the integration of these disciplines, and contribute to the adoption of computer vision by ecologists.

Appendix: Reproducible example of species identification on camera trap images with CPU

In this section, we go through a reproducible example of the entire deep learning workflow, including data preparation, model training, and automatic labeling of new images. We used a subsample of 467 images from the original dataset in the Jura county to allow the training of our model with CPU on a personal computer. We also used 14 images from the original dataset in the Ain county to illustrate prediction.

Training and validation datasets

We first split the dataset of Jura images in two datasets, a dataset for training, and the other one for validation. We use the exifr package to extract metadata from images, get a list of images names and extract the species from these.
Table 4: Species considered, and number of images with these species in them.

| Keywords       | n  |
|----------------|----|
| humain         | 143|
| vehicule       | 135|
| renard         | 58 |
| sangliers      | 33 |
| chasseur       | 17 |
| chien          | 14 |
| lynx           | 13 |
| chevreuil      | 13 |
| chamois        | 12 |
| blaireaux      | 10 |
| chat           | 8  |
| lievre         | 4  |
| fouine         | 1  |
| cavalier       | 1  |

Then we pick 80% of the images for training in each category, the rest being used for validation.

```r
library(exifr)
pix_folder <- '/quotesingle.ts1pix/pixJura/
file_list <- list.files(path = pix_folder,
 recursive = TRUE,
 pattern = "*.JPG",
 full.names = TRUE)

labels <-
 read_exif(file_list) %>%
 as_tibble() %>%
 unnest(Keywords, keep_empty = TRUE) %>% # keep_empty = TRUE keeps pix with no labels (empty pix)
 group_by(SourceFile) %>%
 slice_head() %>% # when several labels in a pix, keep first only
 ungroup() %>%
 mutate(Keywords = as_factor(Keywords)) %>%
 mutate(Keywords = fct_explicit_na(Keywords, "wo_tag")) %>% # when pix has no tag
 select(SourceFile, FileName, Keywords) %>%
 mutate(Keywords = fct_recode(Keywords,
 "chat" = "chat forestier",
 "lievre" = "lièvre",
 "vehicule" = "véhicule",
 "ni" = "Non identifié")) %>%
 filter(!(Keywords %in% c("ni", "wo_tag")))

Then we pick 80% of the images for training in each category, the rest being used for validation.

# training dataset
pix_train <- labels %>%
 select(SourceFile, FileName, Keywords) %>%
 group_by(Keywords) %>%
 filter(between(row_number(), 1, floor(n() * 80/100))) # 80% per category

# validation dataset
pix_valid <- labels %>%
 select(SourceFile, FileName, Keywords) %>%
 group_by(Keywords) %>%
 filter(between(row_number(), floor(n() * 80/100) + 1, n()))
```
Eventually, we store these images in two distinct directories named `train` and `valid`.

```r
# create dir train/ and copy pix there, organised by categories
dir.create('pix/train') # create training directory
for (i in levels(fct_drop(pix_train$Keywords))) dir.create(paste0('pix/train/',i)) # create dir for lab
for (i in 1:nrow(pix_train)){
  file.copy(as.character(pix_train$SourceFile[i]),
            paste0('pix/train/', as.character(pix_train$Keywords[i]))) # copy pix in corresp dir
}
# create dir valid/ and copy pix there, organised by categories.
dir.create('pix/valid') # create validation dir
for (i in levels(fct_drop(pix_train$Keywords))) dir.create(paste0('pix/valid/',i)) # create dir for lab
for (i in 1:nrow(pix_valid)){
  file.copy(as.character(pix_valid$SourceFile[i]),
            paste0('pix/valid/', as.character(pix_valid$Keywords[i]))) # copy pix in corresp dir
}
# delete pictures in valid/ directory for which we did not train the model
if (!is_empty(to_be_deleted)) {
  for (i in 1:length(to_be_deleted)){
    unlink(paste0('pix/valid/', to_be_deleted[i]))
  }
}
```

What is the sample size of these two datasets?

```r
bind_rows("training" = pix_train, "validation" = pix_valid, .id = "dataset") %>%
group_by(dataset) %>%
count(Keywords) %>%
rename(category = Keywords) %>%
kable(caption = "Sample size (n) for the training and validation datasets.") %>%
kable_styling()
```

Transfer learning

We proceed with transfer learning using images from the Jura county (or a subsample more exactly). We first load images and apply standard transformations to improve training (flip, rotate, zoom, rotate, light transform).

```r
dls <- ImageDataLoaders_from_folder(
  path = "pix/",
  train = "train",
  valid = "valid",
  item_tfms = Resize(size = 460),
  bs = 10,
  batch_tfms = list(aug_transforms(size = 224,
                                    min_scale = 0.75), # transformation
                    Normalize_from_stats( imagenet_stats() ))),
  num_workers = 0,
  ImageFile.LOAD_TRUNCATED_IMAGES = TRUE)
```

Then we get the model architecture. For the sake of illustration, we use a resnet18 here, but we used a resnet50 to get the full results presented in the main text.

```r
learn <- cnn_learner(dls = dls,
                      arch = resnet18(),
```
Table 5: Sample size (n) for the training and validation datasets.

| dataset | category | n  |
|---------|----------|----|
| training | humain   | 114|
| training | vehicule | 108|
| training | chamois  | 9  |
| training | blaireaux| 8  |
| training | sangliers| 26 |
| training | renard   | 46 |
| training | chasseur | 13 |
| training | lynx     | 10 |
| training | chien    | 11 |
| training | chat     | 6  |
| training | chevreuil| 10 |
| training | lievre   | 3  |
| validation | humain | 29 |
| validation | vehicule | 27 |
| validation | chamois | 3  |
| validation | blaireaux | 2 |
| validation | sangliers| 7  |
| validation | renard  | 12 |
| validation | chasseur | 4  |
| validation | lynx    | 3  |
| validation | chien   | 3  |
| validation | fouine  | 1  |
| validation | chat    | 2  |
| validation | chevreuil| 3  |
| validation | lievre  | 1  |
| validation | cavalier| 1  |
Now we are ready to train our model. Again, for the sake of illustration, we use only 2 epochs here, but used 20 epochs to get the full results presented in the main text. With all pictures and a resnet50, it took 75 minutes per epoch approximatively on a Mac with a 2.4Ghz processor and 64Go memory, and less than half an hour on a machine with GPU. On this reduced dataset, it took a bit more than a minute per epoch on the same Mac. Note that we save the model after each epoch for later use.

```r
one_cycle <- learn %>%
  fit_one_cycle(2, cbs = SaveModelCallback(every_epoch = TRUE, 
                                          fname = "model"))
```

This yields the following model:

```r
## epoch train_loss valid_loss accuracy error_rate time
## ------ ----------- ----------- --------- ----------- ----- 
## 0 2.447199 0.850911 0.760417 0.239583 01:24 
## 1 1.736201 0.666055 0.781250 0.218750 01:25 
```

We may dig a bit deeper in training performances by loading the best model, here `model_1.pth`, and display some metrics for each species.

```r
learn$load("model_1")
```

This yields the following model:

```r
## Sequential(
##  (0): Sequential(
##    (0): Conv2d(3, 64, kernel_size=(7, 7), stride=(2, 2), padding=(3, 3), bias=False)
##    (1): BatchNorm2d(64, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##    (2): ReLU(inplace=True)
##    (3): MaxPool2d(kernel_size=3, stride=2, padding=1, dilation=1, ceil_mode=False)
##  (4): Sequential(
##    (0): BasicBlock(
##      (conv1): Conv2d(64, 64, kernel_size=(3, 3), stride=(1, 1), padding=(1, 1), bias=False)
##      (bn1): BatchNorm2d(64, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##      (relu): ReLU(inplace=True)
##      (conv2): Conv2d(64, 64, kernel_size=(3, 3), stride=(1, 1), padding=(1, 1), bias=False)
##      (bn2): BatchNorm2d(64, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##    )
##    (1): BasicBlock(
##      (conv1): Conv2d(64, 64, kernel_size=(3, 3), stride=(1, 1), padding=(1, 1), bias=False)
##      (bn1): BatchNorm2d(64, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##      (relu): ReLU(inplace=True)
##      (conv2): Conv2d(64, 64, kernel_size=(3, 3), stride=(1, 1), padding=(1, 1), bias=False)
##      (bn2): BatchNorm2d(64, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##    )
##  )
##  (5): Sequential(
##    (0): BasicBlock(
##      (conv1): Conv2d(64, 128, kernel_size=(3, 3), stride=(2, 2), padding=(1, 1), bias=False)
##      (bn1): BatchNorm2d(128, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##      (relu): ReLU(inplace=True)
##      (conv2): Conv2d(128, 128, kernel_size=(3, 3), stride=(1, 1), padding=(1, 1), bias=False)
##      (bn2): BatchNorm2d(128, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
##    )
##  )
```
## (1): Sequential(
## (0): AdaptiveConcatPool2d(
##     (ap): AdaptiveAvgPool2d(output_size=1)
##     (mp): AdaptiveMaxPool2d(output_size=1)
## )
## (1): Flatten(full=False)
## (2): BatchNorm1d(1024, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
## (3): Dropout(p=0.25, inplace=False)
## (4): Linear(in_features=1024, out_features=512, bias=False)
## (5): ReLU(inplace=True)
## (6): BatchNorm1d(512, eps=1e-05, momentum=0.1, affine=True, track_running_stats=True)
## (7): Dropout(p=0.5, inplace=False)
## (8): Linear(in_features=512, out_features=12, bias=False)
## )
## )

textmlin <- ClassificationInterpretation_from_learner(learn)
textmlin$print_classification_report()

We may extract the categories that get the most confused.

textmlin %>% most_confused()

| V1        | V2     | V3 |
|-----------|--------|----|
| humain    | vehicule | 4  |
| chasseur  | humain  | 3  |
| blaireaux | renard  | 1  |
| blaireaux | sangliers | 1 |
| chat      | renard  | 1  |
| chat      | sangliers | 1 |
| chevreuil | chien   | 1  |
| chevreuil | renard  | 1  |
| chien     | chamois | 1  |
| chien     | sangliers | 1 |
| humain    | chasseur | 1 |
| lievre    | renard  | 1  |
| renard    | blaireaux | 1 |
| renard    | sangliers | 1 |
| sangliers | renard  | 1  |
| vehicule  | humain  | 1  |

**Transferability**

In this section, we show how to use our freshly trained model to label images that were taken in another study site in the Ain county, and not used to train our model. First, we get the path to the images.

```r
tfls <- list.files(path = "pix/pixAin",
                   full.names = TRUE,
                   recursive = TRUE)
```

Then we carry out prediction, and compare to the truth.

```r
predicted <- character(3)
categories <- interp$vocab %>%
     str_replace_all("[[:punct:]]", " ") %>%
     str_trim() %>%
     str_split(" ") %>%
```
Table 6: Comparison of the predictions vs. ground truth.

| truth     | prediction |
|-----------|------------|
| lynx      | chevreuil  |
| roe deer  | chamois    |
| wild boar | sangliers  |

```
unlist()
for (i in 1:length(fls)){
  result <- learn %>% predict(fls[i]) # make prediction
  result[[3]] %>%
    str_extract("\\d+") %>%
    as.integer() -> index # extract relevant info
  predicted[i] <- categories[index + 1] # match it with categories
}
data.frame(truth = c("lynx", "roe deer", "wild boar"),
            prediction = predicted) %>%
kable(caption = "Comparison of the predictions vs. ground truth.") %>%
kable_styling()
```

Session information

````
## R version 4.1.0 (2021-05-18)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Catalina 10.15.7
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/4.1/Resources/lib/libRblas.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.1/Resources/lib/libRlapack.dylib
##
## locale:
## [1] fr_FR.UTF-8/fr_FR.UTF-8/fr_FR.UTF-8/C/fr_FR.UTF-8/fr_FR.UTF-8/fr_FR.UTF-8
##
## attached base packages:
## [1] stats graphics grDevices utils datasets methods base
##
## other attached packages:
## [1] exifr_0.3.2 unmarked_1.1.9006 lattice_0.20-44
## [4] janitor_2.1.0 highcharter_0.8.2 fastai_2.0.9
## [7] ggtext_0.1.1 wesanderson_0.3.6 kableExtra_1.3.4
## [10] stringi_1.7.3 lubridate_1.7.10 cowplot_1.1.1
## [13] sf_1.0-2forcats_0.5.1 stringr_1.4.0
## [16] dplyr_1.0.7 purrr_0.3.4.9000 readr_2.0.0
## [19] tidyr_1.1.3 tibble_3.1.3 ggplot2_3.3.5
## [22] tidyverse_1.3.1
##
## loaded via a namespace (and not attached):
## [1] minqa_1.2.4 colorspace_2.0-2 gg-gsignif_0.6.2
## [4] rio_0.5.27 ellipsis_0.3.2 class_7.3-19
## [7] snakecase_0.11.0 markdown_1.1 fs_1.5.0
## [10] gridtext_0.1.4 rstudioapi_0.13 proxy_0.4-26
```

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