Biodiversity in National Parks

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1: Introduction

The "biodiversity.ipynb" contains all the information and code for the project

The "biodiversity.csv" contains the cleaned and merged data from "observations.csv" and "species_info.csv", which was curated by National Parks Service (https://www.nps.gov/index.htm).

The objective of this project was to analyze the biodiversity data in order to better understand the level of endangerment of certain species in various parks along with the frequency of their sightings.

Endangered species are living organisms, both plants and animals, that face a high risk of extinction.

These species encounter significant threats caused by diverse factors such as habitat loss, climate change, pollution, poaching, and the introduction of invasive species.

The declining population numbers of these species raise serious concerns about the delicate equilibrium of biodiversity on our planet.

Biodiversity encompasses the remarkable array of life forms, ranging from microorganisms to plants, animals, and entire ecosystems.

It encompasses the intricate network of interactions between different species and their surrounding environment. Therefore, it plays a critical role in sustaining the health and functionality of ecosystems, which, in turn, supports life on Earth.

The conservation of biodiversity is crucial for ensuring the continued well-being of our planet and the future of all living beings.

2: Modules

```
import re
import numpy as np
import pandas as pd
import seaborn as sns
import matplotlib.pyplot as plt
import matplotlib.ticker as ticker
import plotly.express as px
import plotly.graph_objects as go
from scipy.stats import entropy
from scipy.stats import thenepexact
from scipy.stats import barnand_exact
from scipy.stats import contingency
```

3: Functions

```
In [2]: # function to get dataframe name: Useful only for missing data check function
               name =[x for x in globals() if globals()[x] is df][0]
            function that checks if any data is missing, identifies the data type, and displays the first 10 row indices where missing data is found
          def missing_data_check(data):
    print(f'Missing data check for "{get_df_name(data)}" dataframe:', "\n")
               nan_values = data.isna().any().any()
               nan_values = data.isna().any().any()
missing_values = data.empty
empty_string_values = (data == "").any().any()
print(f"NaN values exist: {nan_values}")
print(f"Missing values exist: {missing_values}")
               print(f"Empty string values exist: {empty_string_values}")
               if nan_values or missing_values or empty_string_values:
    print("\n" + "Rows with missing or empty values:")
                    if nan values:
                         nan_rows = data[data.isna().any(axis=1)]
                         print("NaN rows
                         print(nan_rows.index.tolist()[:10], "\n")
                    if missing_values:
                         missing_rows = data[data.empty]
print("Missing rows:")
                         print(missing_rows.index.tolist()[:10], "\n")
                    if empty string values:
                         empty_string_rows = data[data == ""]
                         print("Empty string rows:
                         print(empty_string_rows.index.tolist()[:10], "\n")
               else:
                    print("\n" + "No missing or empty values found.", "\n")
```

```
In [3]: # function that calculates Chi2, Fisher's, Barnard's and Boschloo's exact tests and stores the data into a "exact_tests_results_df" dataframe
# Boschloo's exact test is not considered in this analysis due to its use for ordinal categories
# example: contingency_table = [[584, 120], [288, 28]]

def calculate_exact_tests(contingency_table):
    chi2_result = chi2_contingency(contingency_table)
    fisher_result = fisher_exact(contingency_table)
    barnard_result = barnard_exact(contingency_table)
# boschloo_result = boschloo_exact(contingency_table)

exact_tests_results_df = pd.DataFrame(columns=["Test", "Statistic", "p-value"])
    exact_tests_results_df.loc[0] = ["Chi2", chi2_result.statistic, chi2_result.pvalue]
    exact_tests_results_df.loc[1] = ["Fisher", fisher_result.statistic, fisher_result.pvalue]
    exact_tests_results_df.loc[2] = ["Barnard", barnard_result.statistic, barnard_result.pvalue]

# exact_tests_results_df.loc[3] = ["Boschloo", boschloo_result.statistic, boschloo_result.pvalue]

return exact_tests_results_df.
```

4: Data Cleaning

The primary focus was placed on cleaning the data, with the objective of obtaining a singular dataset that is entirely pristine and suitable for analysis.

Initially, a thorough examination and refinement process was conducted on both files individually. Subsequently, these files were merged, and an additional round of filtering was performed, ensuring the creation of a **single, definitive ".csv" file**.

4.0: Personal Preference Options

```
In [4]: # personal preference settings
pd.set_option("display.max_columns", None)
pd.set_option("display.max_ros", None)
pd.set_option("display.max_colwidth", None)
pd.set_option("expand_frame_repr", False)

# Load the csv files into DataFrames
observations = pd.read_csv("observations.csv")
species = pd.read_csv("species_info.csv")
```

4.1: observations.csv

```
observations.info()
        <class 'pandas.core.frame.DataFrame'
        RangeIndex: 23296 entries, 0 to 23295
       Data columns (total 3 columns):

# Column Non-Null Count Dtype
         0 scientific_name 23296 non-null object
        1 park_name 23296 non-null object
2 observations 23296 non-null int64
        dtypes: int64(1), object(2)
memory usage: 546.1+ KB
In [6]: # a quick glance into the data
         observations.head()
                    scientific_name
                 Vicia benghalensis Great Smoky Mountains National Park
         0
                                                                                    68
         1 Neovison vison Great Smoky Mountains National Park
                                                                                   77
                  Prunus subcordata
                                                  Yosemite National Park
         3 Abutilon theophrasti Bryce National Park
                                                                                    84
          4 Githopsis specularioides Great Smoky Mountains National Park
                                                                                    85
In [7]: missing_data_check(observations)
       Missing data check for "observations" dataframe:
        NaN values exist: False
        Missing values exist: False
        Empty string values exist: False
        No missing or empty values found.
         observations.park_name.nunique()
Out[8]: 4
In [9]: # unique park places names
          observations.park_name.unique()
Out[9]: array(['Great Smoky Mountains National Park', 'Yosemite National Park', 'Bryce National Park', 'Yellowstone National Park'], dtype=object)
```

The "observations.csv" file contains three columns:

observations.scientific_name.nunique()

Scientific Name

Out[10]: 5541

Latin name of species. There are 5541 distinct species in this data.

Park Name

There are four different parks in the observations file:

- 1. Great Smoky Mountains National Park
- 2. Yosemite National Park
- 3. Bryce National Park
- 4. Yellowstone National Park
- Observations

There are no null/NaN/missing values.

4.2: species.csv

```
In [11]: # general file structure
          species.info()
        <class 'pandas.core.frame.DataFrame'>
        RangeIndex: 5824 entries, 0 to 5823
        Data columns (total 4 columns):
                           Non-Null Count Dtype
        # Column
         0 category
                                   5824 non-null
                                                    object
         1 scientific_name
                                   5824 non-null
                                                    object
         2 common_names
                                   5824 non-null
                                                    obiect
             conservation_status 191 non-null
                                                    object
        dtypes: object(4)
        memory usage: 182.1+ KB
In [12]: # a quick glance into the data
         species.head()
            category
                                   scientific name
                                                                                          common_names conservation_status
         0 Mammal Clethrionomys gapperi gapperi
                                                                                  Gapper's Red-Backed Vole
                                   Bos bison
         1 Mammal
                                                                                      American Bison, Bison
                                                                                                                        NaN
          2 Mammal
                                        Bos taurus Aurochs, Aurochs, Domestic Cattle (Feral), Domesticated Cattle
                                                                                                                         NaN
                                      Ovis aries
         3 Mammal
                                                     Domestic Sheep, Mouflon, Red Sheep, Sheep (Feral)
                                                                                                                        NaN
          4 Mammal
                                    Cervus elaphus
                                                                                             Wapiti Or Elk
                                                                                                                         NaN
In [13]: missing_data_check(species)
        Missing data check for "species" dataframe:
        NaN values exist: True
        Missing values exist: False
        Empty string values exist: False
        Rows with missing or empty values:
        [0, 1, 2, 3, 4, 5, 6, 10, 11, 12]
In [14]: # where are NaN values? conservation_status!
         species.isna().sum()
Out[14]: category
                                     0
          scientific_name
                                     0
         common_names
          conservation_status
                                  5633
         dtype: int64
In [15]: # number of unique conservation status groups
         species.conservation_status.nunique()
Out[15]: 4
In [16]: # unique conservations status group names
         species.conservation_status.unique()
Out[16]: array([nan, 'Species of Concern', 'Endangered', 'Threatened',
                  'In Recovery'], dtype=object)
In [17]: # number of unique species in the data
         # !WRONG! Although there is a correct number of distinct species in the "species.csv" file,
# it should be noted that there is a discrepancy based on the output of "species.info()"!
         species.scientific_name.nunique()
Out[17]: 5541
In [18]: # there is a discrepancy in the number od species in both datasets (5541 in "observations.csv" / 5824 in "species.csv"), # this can be observed with the "species.info()" function as well!
         len(species.scientific_name)
Out[18]: 5824
In [19]: # more depth for the "NaN" values
         display(species.conservation_status.value_counts())
        conservation_status
        Species of Concern 161
        Endangered
                         10
        Threatened
        In Recovery
        Name: count, dtvpe: int64
In [20]: # percentage of "nan" values for conservation status print("\n" + f'"NaN" values in "conservation_status" constitute ' + str(round(5633 / 5824 * 100, 2)) + "% of the whole dataset!")
        "NaN" values in "conservation_status" constitute 96.72% of the whole dataset!
In [21]: # Discarding over 95 % of the data isn't favorable.
                 "NaN" conservation status has been replaced with "Least concern," enabling its inclusion in the analysis.
         species["conservation_status"].fillna("Least concern", inplace=True)
In [22]: # number of unique categories
         species.category.nunique()
Out[22]: 7
In [23]: # unique categories
         species.category.unique()
Out[23]: array(['Mammal', 'Bird', 'Reptile', 'Amphibian', 'Fish', 'Vascular Plant',
                  'Nonvascular Plant'], dtype=object)
```

```
In [24]: # a quick glance at the common names
          species["common_names"].head()
                                                           Gapper's Red-Backed Vole
Out[24]: 0
                                                               American Bison, Bison
               Aurochs, Aurochs, Domestic Cattle (Feral), Domesticated Cattle
                              Domestic Sheep, Mouflon, Red Sheep, Sheep (Feral)
                                                                       Wapiti Or Elk
          Name: common_names, dtype: object
In [25]: # by observing the common names from the "species.csv", we can see the usual delimiters in these examples
          species["common_names"].iloc[[2, 4, 16, 232, 233, 750, 997, 2295, 2554]]
                   Aurochs, Aurochs, Domestic Cattle (Feral), Domesticated Cattle
Out[25]: 2
                                                               Wapiti Or Elk
Panther (Mountain Lion)
          16
          232
233
                                                    Baltimore Oriole, Northern Oriole
Orchard Oriole
          750
                                                                          Curtis? Aster
                                                                   Venus? Looking-Glass
           997
                                                      A Bluegrass, Bluegrass
A Bramble, Truculent Blackberry
           2295
           2554
          Name: common_names, dtype: object
In [26]: # regular expression that matches ",", letters "Or" together and not in a word (so not 'abort' and such) and "(" regex = r",|\b0r\b|\("
          # clean the common names column and keep only the first common name
species["common_names"] = species["common_names"].apply(lambda x: re.split(regex, x)[0].strip())
          # remove the capital letter "A" with a whitespace after it
species["common_names"] = species["common_names"].apply(lambda x: re.sub(r"A\s", "", x))
           # substitute "?" with whitespace
          species["common\_names"] = species["common\_names"].apply(lambda \ x: \ re.sub(r"\?", \ "", \ x))
In [27]: # check if the correction works
           species["common_names"].iloc[[2, 4, 16, 232, 233, 750, 997, 2295, 2554]]
Out[27]: 2
                                Aurochs
                                  Wapiti
                                Panther
          232
                     Baltimore Oriole
                      Orchard Oriole
                           Curtis Aster
           750
           997
                   Venus Looking-Glass
          2295
                            Bluegrass
          Name: common_names, dtype: object
In [28]: # initially, the regex method removed entire common names for some species, which is the reason for conducting this check and for making the function in the first place.
          # Empty string (" ") isn't the same as NaN, so ".isna()" isn't viable here...
missing_data_check(species)
        Missing data check for "species" dataframe:
         NaN values exist: False
         Missing values exist: False
        Empty string values exist: False
        No missing or empty values found.
In [29]: # number of duplicate scientific names in the data
          duplicated_scientific_names = species["scientific_name"].duplicated()
          print(duplicated_scientific_names.value_counts(), "\n")
          # number of duplicate common names in the data.
# Not relevant since I'm planning to keep only the first common
duplicated_common_names = species["common_names"].duplicated()
                                                                                on name to make the dataset cleaner!
          print(duplicated_common_names.value_counts())
         scientific_name
         False 5541
         True
                    283
         Name: count, dtype: int64
         common_names
         False 5291
                   533
         Name: count, dtype: int64
In [30]: # store the duplicates to better understand how to clean the
          duplicates = species[species["scientific_name"].duplicated()]
           # a quick glance into the duplicates
          duplicates.head()
Out[30]:
                                  scientific name common names conservation status
                 category
           3017 Mammal
                                   Cervus elaphus Rocky Mountain Elk
          3019 Mammal Odocoileus virginianus White-Tailed Deer
                                                                             Least concern
           3020 Mammal
                                                            Gray Wolf
                                                                              In Recovery
                                      Canis lupus
                                                   Cougar
          3022 Mammal
                            Puma concolor
                                                                        Least concern
           3025 Mammal
                                 Lutra canadensis
                                                          River Otter
                                                                             Least concern
In [31]: # seeing if all duplicates are the same, chosen at random from the .head() display
          display(species[species["scientific_name"] == "Cervus elaphus"])
display(species[species["scientific_name"] == "Lutra canadensis"])
          # there is a discrepancy in the conservation status of some (how many?) duplicates
display(species[species["scientific_name"] == "Canis lupus"])
                category scientific_name common_names conservation_status
            4 Mammal Cervus elaphus
                                                       Wapiti
                                                                     Least concern
         3017 Mammal Cervus elaphus Rocky Mountain Elk
                category scientific_name
                                              common names conservation status
           20 Mammal Lutra canadensis Northern River Otter
                                                                       Least concern
                                           River Otter
         3025 Mammal Lutra canadensis
                                                                      Least concern
```

```
8 Mammal
                              Canis lupus
                                                Gray Wolf
                                                                   Endangered
        3020 Mammal Canis lupus Gray Wolf
                                                                  In Recovery
         4448 Mammal
                             Canis lupus
                                                Gray Wolf
                                                                   Endangered
In [32]: # find duplicates based on scientific name
          dups = species[species.duplicated(subset="scientific_name", keep=False)]
          # find duplicates with different conservation status.
# "~" is a negation operator which ensures that only duplicates with different conservation statuses are stored
          \label{thm:conservation} $$\operatorname{dups\_duplicated(subset="scientific\_name", keep=False)} \& \operatorname{\neg dups\_duplicated(subset="conservation\_status", keep=False)}]$
          display(dups_with_diff_conservation_status)
               category
                              scientific_name common_names conservation_status
                                                                        In Recovery
        3020 Mammal
                                  Canis lupus
                                                     Grav Wolf
        3283 Fish Oncorhynchus mykiss Rainbow Trout
                                                                        Threatened
In [33]: species[species["scientific_name"] == "Oncorhynchus mykiss"]
                 category
                                scientific_name common_names conservation_status
            560
                     Fish Oncorhynchus mykiss Rainbow Trout
                                                                        Least concern
          3283
                    Fish Oncorhynchus mykiss Rainbow Trout
                                                                       Threatened
In [34]: # since each of the first duplicate species has the correct conservation status, this should yield a clean dataset.
          species.drop\_duplicates(subset=["scientific\_name"], \ keep="first", \ inplace=True)
In [35]: # double-check
          display(species[species["scientific_name"] == "Canis lupus"])
display(species[species["scientific_name"] == "Oncorhynchus mykiss"])
            category scientific_name common_names conservation_status
        8 Mammal
                          Canis lupus
                                             Gray Wolf
                                                               Endangered
              category
                             scientific_name common_names conservation_status
                 Fish Oncorhynchus mykiss Rainbow Trout
        560
                                                                    Least concern
In [36]: # verify the number of distinct species in each dataset; Checks out!
          print(observations.scientific_name.nunique())
print(species.scientific_name.nunique())
          print(len(species.scientific_name))
        5541
         5541
```

The "species.csv" file contains four columns:

Category

5541

There are seven distinct categories in the data:

category scientific_name common_names conservation_status

- 1. Mammal
- 2. Bird
- 3. Reptile
- 4. Amphibian
- 5. Fish
- 6. Vascular Plant
- 7. Nonvascular Plant

• Scientific Name

Apperas to be the same naming as in "observations.csv", although with a different number of distinct species (before cleaning)

Common Names

One or more common names for different species. Only the first common name is preserved after cleaning.

• Conservation Status

Refers to the assessment of the risk level faced by a species or ecosystem, indicating the extent to which it is threatened or protected based on scientific evaluation and criteria. The dataset includes **four distinct conservation status groups**, along with the majority classified as "NaN" (before cleaning):

- 1. Species of Concern
- 2. Endangered
- 3. Threatened
- 4. In Recovery
- 5. "NaN"

According to Wikipedia (https://en.wikipedia.org/wiki/Conservation_status), species are classified by the IUCN Red List into nine groups:

- 1. Extinct (EX) No known living individuals.
- 2. Extinct in the wild (EW) Known only to survive in captivity, or as a naturalized population outside its historic range.
- 3. Critically Endangered (CR) Highest risk of extinction in the wild.
- 4. **Endangered (EN)** Higher risk of extinction in the wild.
- 5. Vulnerable (VU) High risk of extinction in the wild.
- 6. Near Threaened (NT) Likely to become endangered in the near future.
- 7. Conservation Dependent (CD) Low risk; is conserved to prevent being near threatened, certain events may lead it to being a higher risk level.
- 8. Least concern (LC) Very Low risk; does not qualify for a higher risk category and not likely to be threatened in the near future. Widespread and abundant taxa are included in this category.

9. Data deficient (DD) - Not enough data to make an assessment of its risk of extinction.

10. Not evaluated (NE) – Has not yet been evaluated against the criteria.

The "NaN" group is replaced with "Least concern" in the "species.csv".

There is a discrepancy between the number of distinct species and the conservation status of some of them.

U.S. Fish & Wildlife Service categorizes Canis lupus as "Endangered" (https://www.fws.gov/initiative/protecting-wildlife/gray-wolf-recovery-news-and-updates).

The species Oncorhynchus mykiss is not listed (https://ecos.fws.gov/ecp/species/757).

However, in certain states, it is classified as a "Pest Species." Therefore, considering the existing distributions, the "Least Concern" group is appropriate.

Duplicates have been identified and subsequently removed, thereby preserving the accurate conservation status of each species based on its initial occurrence and/or through external verification.

4.3: Merged Data

biodiversity_dirty.head()

```
In [37]: # merge column will be "scientific_name"
display(observations.head())
          display(species.head())
          # merge the data into a single dataframe
          biodiversity_dirty = pd.merge(observations, species, on="scientific_name")
                  scientific name
                                                          park name observations
        0
                Vicia benghalensis Great Smoky Mountains National Park
           Neovison vison Great Smoky Mountains National Park
        2
                Prunus subcordata
                                               Yosemite National Park
                                                                               138
              Abutilon theophrasti
                                                  Bryce National Park
        4 Githopsis specularioides Great Smoky Mountains National Park
                                                                                85
                                  scientific_name
                                                           common_names conservation_status
            category
        0 Mammal Clethrionomys gapperi gapperi Gapper's Red-Backed Vole
                                                                                   Least concern
                                       Bos bison American Bison
                                                                  Aurochs
        2 Mammal
                                        Bos taurus
                                                                                   Least concern
                       Ovis aries Domestic Sheep
        3 Mammal
                                                                                  Least concern
         4 Mammal
                                 Cervus elaphus
                                                                   Wapiti
                                                                                  Least concern
In [38]: # check if the numbers match; They do!
          print(len(observations))
          print(len(biodiversity_dirty))
          print(observations.scientific_name.nunique())
          print(species.scientific_name.nunique())
          print(biodiversity_dirty.scientific_name.nunique(), "\n")
          # the expected number of rows could be four times the number of distinct species in the data if each species is observed in each park!
print(f"Possible expected number of rows: {biodiversity_dirty.scientific_name.nunique() * 4}")
print(f"Difference: " + str(23296 - 22164))
        23296
        23296
        5541
5541
        5541
        Possible expected number of rows: 22164
        Difference: 1132
In [39]: #
          biodiversity_dirty.info()
        <class 'pandas.core.frame.DataFrame'>
        RangeIndex: 23296 entries, 0 to 23295
        Data columns (total 6 columns):
                                    Non-Null Count Dtype
         # Column
         0 scientific_name
                                    23296 non-null object
             park_name
                                   23296 non-null object
             observations
                                   23296 non-null int64
            category
common_names
                                    23296 non-null object
23296 non-null object
            conservation_status 23296 non-null object
        dtypes: int64(1), object(5)
         memory usage: 1.1+ MB
In [40]: missing_data_check(biodiversity_dirty)
        Missing data check for "biodiversity_dirty" dataframe:
        NaN values exist: False
        Missing values exist: False
        Empty string values exist: False
        No missing or empty values found.
```

```
0 Vicia benghalensis Great Smoky Mountains National Park
                                                                               68 Vascular Plant
                                                                                                      Purple Vetch
                                                                                                                          Least concern
                                          Yosemite National Park
                                                                             148 Vascular Plant
          1 Vicia benghalensis
                                                                                                     Purple Vetch
                                                                                                                          Least concern
           2 Vicia benghalensis
                                           Yellowstone National Park
                                                                             247 Vascular Plant
                                                                                                      Purple Vetch
                                                                                                                          Least concern
          3 Vicia benghalensis
                                                 Bryce National Park
                                                                             104 Vascular Plant
                                                                                                     Purple Vetch
                                                                                                                          Least concern
                Neovison vison Great Smoky Mountains National Park
                                                                               77
                                                                                       Mammal
                                                                                                   American Mink
                                                                                                                          Least concern
In [42]: # strip the "National Park" from park names
          biodiversity_dirty["park_name"] = biodiversity_dirty["park_name"].map(lambda x: x.rstrip("National Park"))
          biodiversity_dirty.head()
                scientific_name
                                           park_name observations
                                                                          category common_names conservation_status
           0 Vicia benghalensis Great Smoky Mountains
                                                                  68 Vascular Plant
                                                                                         Purple Vetch
                                                                                                             Least concern
           1 Vicia benghalensis
                                              Yosemite
                                                                 148 Vascular Plant
                                                                                         Purple Vetch
                                                                                                             Least concern
           2 Vicia benghalensis
                                            Yellowstone
                                                                 247 Vascular Plant
                                                                                         Purple Vetch
                                                                                                             Least concern
           3 Vicia benghalensis
                                                 Bryce
                                                                 104 Vascular Plant
                                                                                         Purple Vetch
                                                                                                             Least concern
                Neovison vison Great Smoky Mountains
                                                                           Mammal
                                                                                      American Mink
                                                                  77
                                                                                                             Least concern
In [43]: # check the dataframe for random species; Notice that observations can be summed up to reduce the number of rows in the dataset!
display(biodiversity_dirty[biodiversity_dirty["scientific_name"] == "Helianthus annuus"])
          display(biodiversity_dirty[biodiversity_dirty["scientific_name"] == "Plantago lanceolata"])
                scientific name
                                            park_name observations
                                                                           category
                                                                                       common names conservation status
         268 Helianthus annuus
                                            Yellowstone
                                                                  265 Vascular Plant Common Sunflower
                                                                                                                Least concern
         269
              Helianthus annuus
                                                                 140 Vascular Plant Common Sunflower
                                                                                                                Least concern
         270 Helianthus annuus
                                            Yellowstone
                                                                  235 Vascular Plant Common Sunflower
                                                                                                                Least concern
        271 Helianthus annuus
                                                 Bryce
                                                                  118 Vascular Plant Common Sunflower
                                                                                                                Least concern
         272 Helianthus annuus
                                                                  123 Vascular Plant Common Sunflower
                                               Yosemite
                                                                                                                Least concern
        273 Helianthus annuus
                                              Yosemite
                                                                  169 Vascular Plant Common Sunflower
                                                                                                                Least concern
         274 Helianthus annuus Great Smoky Mountains
                                                                   44 Vascular Plant Common Sunflower
                                                                                                                Least concern
                                                                   54 Vascular Plant Common Sunflower
         275 Helianthus annuus Great Smoky Mountains
                                                                                                                Least concern
                    scientific_name
                                               park_name observations
                                                                              category common_names conservation_status
         20900 Plantago lanceolata Great Smoky Mountains
                                                                      51 Vascular Plant
                                                                                          English Plantain
                                                                                                                 Least concern
         20901 Plantago lanceolata
                                                  Yosemite
                                                                     119 Vascular Plant
                                                                                          English Plantain
                                                                                                                 Least concern
         20902 Plantago lanceolata
                                                                     119 Vascular Plant
                                                                                          English Plantain
                                                     Bryce
                                                                                                                 Least concern
                                                                     136 Vascular Plant
         20903 Plantago lanceolata
                                                 Yosemite
                                                                                          English Plantain
                                                                                                                Least concern
         20904 Plantago lanceolata Great Smoky Mountains
                                                                      79 Vascular Plant
                                                                                          English Plantain
                                                                                                                 Least concern
         20905 Plantago lanceolata
                                                     Bryce
                                                                     137 Vascular Plant
                                                                                          English Plantain
                                                                                                                Least concern
         20906 Plantago lanceolata
                                               Yellowstone
                                                                     260 Vascular Plant
                                                                                          English Plantain
                                                                                                                 Least concern
         20907 Plantago lanceolata
                                              Yellowstone
                                                                     264 Vascular Plant English Plantain
                                                                                                                Least concern
In [44]: # count the number of times a species occurs in the data
          scientific_name_counts = biodiversity_dirty["scientific_name"].value_counts()
           # sort the counts from smallest to largest
           sorted_counts = scientific_name_counts.sort_values()
           # display least and most occuring species
          print(sorted_counts[1:6], "\n")
          print(sorted_counts[-6:-1])
         scientific_name
         Limosa fedoa
         Juncus uncialis
         Thaspium barbinode
         Polystichum scopulinum
         Comandra umbellata
         Name: count, dtype: int64
         scientific name
         Castor canadensis
                                    12
         Streptopelia decaocto
         Holcus lanatus
                                    12
         Hypochaeris radicata
                                    12
         Puma concolor
                                    12
         Name: count, dtype: int64
In [45]: # check the observations for Least and most occurrences
          display(biodiversity_dirty[biodiversity_dirty["scientific_name"] == "Limosa fedoa"])
display(biodiversity_dirty[biodiversity_dirty["scientific_name"] == "Holcus lanatus"])
                scientific_name
                                           park_name observations category common_names conservation_status
         1388
                  Limosa fedoa
                                           Yellowstone
                                                                 266
                                                                           Bird
                                                                                 Marbled Godwit
                                                                                                    Species of Concern
         1389
                  Limosa fedoa
                                                                 131
                                                                                 Marbled Godwit
                                                                                                    Species of Concern
                                              Yosemite
                                                                           Bird
         1390
                  Limosa fedoa Great Smoky Mountains
                                                                  82
                                                                           Bird
                                                                                 Marbled Godwit
                                                                                                    Species of Concern
         1391
                  Limosa fedoa
                                                 Bryce
                                                                  96
                                                                           Bird
                                                                                Marbled Godwit
                                                                                                   Species of Concern
```

category common_names conservation_status

Out[41]:

scientific name

park name observations

```
11634
                 Holcus lanatus
                                                                 83 Vascular Plant Common Velvet Grass
                                                Bryce
                                                                                                               Least concern
                                                                179 Vascular Plant Common Velvet Grass
        11635
                 Holcus lanatus
                                             Yosemite
                                                                                                              Least concern
        11636
                 Holcus lanatus
                                             Yosemite
                                                                138 Vascular Plant Common Velvet Grass
                                                                                                               Least concern
        11637
                 Holcus lanatus
                                           Bryce
                                                                117 Vascular Plant Common Velvet Grass
                                                                                                               Least concern
                                                                 96 Vascular Plant Common Velvet Grass
        11638
                 Holcus lanatus
                                                Bryce
                                                                                                               Least concern
        11639
                 Holcus lanatus Great Smoky Mountains
                                                                 65 Vascular Plant Common Velvet Grass
                                                                                                               Least concern
                                           Yellowstone
                                                                256 Vascular Plant Common Velvet Grass
        11640
                 Holcus lanatus
                                                                                                               Least concern
        11641
                 Holcus lanatus Great Smoky Mountains
                                                                77 Vascular Plant Common Velvet Grass
                                                                                                               Least concern
                 Holcus lanatus Great Smoky Mountains
                                                                 74 Vascular Plant Common Velvet Grass
        11642
                                                                                                               Least concern
        11643
                 Holcus lanatus
                                          Yellowstone
                                                                287 Vascular Plant Common Velvet Grass
                                                                                                               Least concern
In [46]: # check if there is any species that isn't observed in every park in the data
            a list of all park names
          all_park_names = biodiversity_dirty["park_name"].unique()
          # check which species aren't observed in every park
          not_in_every_park
          for species in biodiversity_dirty["scientific_name"].unique():
               species_parks = biodiversity_dirty[biodiversity_dirty["scientific_name"] == species]["park_name"].unique()
              if len(species_parks) != len(all_park_names):
                  not_in_every_park.append(species)
            print the list of species not found in every park or a message if all species are observed in every park
         if not_in_every_park:
              print("The following species are not observed in every park:")
              for species in not_in_every_park:
                  print(species)
          else:
              print("All species are observed in every park!")
        All species are observed in every park!
            group by park name and sum observations for each scientific name
         biodiversity = biodiversity_dirty.groupby(["park_name", "scientific_name", "category", "common_names", "conservation_status"])["observations"].sum().reset_index()
           # inspect the cleaned DataFrame
         display(biodiversity.head())
         # double-check the observations for random species
display(biodiversity[biodiversity["scientific_name"] == "Helianthus annuus"])
display(biodiversity[biodiversity["scientific_name"] == "Holcus lanatus"])
                          scientific_name
                                                                    common_names conservation_status observations
           park_name
                                                  category
        0
                              Abies bifolia
                                              Vascular Plant Rocky Mountain Alpine Fir
                                                                                                                   109
                 Bryce
                                                                                            Least concern
                                              Vascular Plant
                                                                          Balsam Fir
                                                                                        Least concern
                                                                                                                   83
                 Bryce
                           Abies concolor
        2
                 Brvce
                              Ahies fraseri
                                              Vascular Plant
                                                                           Fraser Fir Species of Concern
                                                                                                                   109
                 Bryce Abietinella abietina Nonvascular Plant
                                                                    Abietinella Moss
                                                                                           Least concern
                                              Vascular Plant Wyoming Sand Verbena Species of Concern
                 Brvce Abronia ammophila
                                                                                                                   92
                           park_name scientific_name
                                                                       common_names conservation_status observations
                                                             category
                                Bryce Helianthus annuus Vascular Plant Common Sunflower
         2368
                                                                                                 Least concern
                                                                                                                       258
         7909
               Great Smoky Mountains Helianthus annuus Vascular Plant Common Sunflower
                                                                                                 Least concern
                                                                                                                        98
                           Yellowstone Helianthus annuus Vascular Plant Common Sunflower
        13450
                                                                                                 Least concern
                                                                                                                        500
        18991
                             Yosemite Helianthus annuus Vascular Plant Common Sunflower
                                                                                                 Least concern
                                                                                                                       292
                           park_name scientific_name
                                                                         common_names conservation_status observations
                                                           category
         2452
                                Bryce Holcus lanatus Vascular Plant Common Velvet Grass
                                                                                                 Least concern
                                                                                                                       296
         7993 Great Smoky Mountains Holcus lanatus Vascular Plant Common Velvet Grass
                                                                                                 Least concern
                                                                                                                       216
        13534
                          Yellowstone Holcus lanatus Vascular Plant Common Velvet Grass
                                                                                                                       805
                                                                                                 Least concern
        19075
                             Yosemite Holcus lanatus Vascular Plant Common Velvet Grass
                                                                                                 Least concern
                                                                                                                       463
In [48]: # general datafro
                             ne structure
         biodiversity.info()
        <class 'pandas.core.frame.DataFrame'
        RangeIndex: 22164 entries, 0 to 22163
        Data columns (total 6 columns):
                                    Non-Null Count Dtype
             Column
             park name
                                    22164 non-null object
              scientific_name
                                    22164 non-null
                                    22164 non-null object
             category
                                    22164 non-null object
             conservation_status 22164 non-null
                                                     object
             observations
                                    22164 non-null int64
        dtypes: int64(1), object(5)
        memory usage: 1.0+ ME
In [49]: # double-check species number
         print(biodiversity["scientific name"].nunique())
          # expected number of rows; It is confirmed in the cells above that each species is observed in each park,
         # therefore the expected number of rows is four times the number of species, after summing the observations for each park! print(biodiversity["scientific_name"].nunique() * 4)
          # double-check row number without multiple observations for the same park; Checks out!
         print(len(biodiversity))
        5541
        22164
        22164
```

scientific name

Holcus lanatus

Holcus lanatus

In [50]: missing_data_check(biodiversity)

11632

11633

park name observations

Yellowstone

Yosemite

category

262 Vascular Plant Common Velvet Grass

146 Vascular Plant Common Velvet Grass

common names conservation status

Least concern

Least concern

```
Empty string values exist: False

No missing or empty values found.

In [51]: # export the data into a "biodiversity.csv" file
```

The "biodiversity.csv" file contains six columns, cleaned and merged from "observations.csv" and "species.csv":

Scientific Name

NaN values exist: False Missing values exist: False

There are 5541 distinct species in this dataset with corresponding Latin names.

Park Name

There are four distinct parks in this dataset:

biodiversity.to_csv("biodiversity.csv", index=False)

- 1. Great Smoky Mountains National Park
- 2. Yosemite National Park

Missing data check for "biodiversity" dataframe:

- 3. Bryce National Park
- 4. Yellowstone National Park
- Observations

The count of species sightings per park for each individual species. Each species appeares in each park.

Category

There are seven distinct categories in the data:

- 1. Mammal
- 2. Bird
- 3. Reptile
- 4. Amphibian
- 5. Fish
- 6. Vascular Plant
- 7. Nonvascular Plant
- Common Names

Well-known names commonly recognized by the general population.

Conservation Status

Refers to the assessment of the risk level faced by a species or ecosystem, indicating the extent to which it is threatened or protected based on scientific evaluation and criteria.

The dataset includes **five distinct conservation status groups**:

- 1. Species of Concern
- 2. Endangered
- 3. Threatened
- 4. In Recovery
- 5. Least Concern

Title Figure 1:

Biodiversity in Focus: Exploring Species Endangerment and Sightings across Four American National Parks

```
In [52]: # national park names, states and coordinates
                       "Location": ["Bryce Canyon, Utah", "Great Smoky Mountains, Tennessee", "Yosemite, California", "Yellowstone, Wyoming"], "Latitude": [37.5930, 35.6131, 37.8651, 44.4279],
                       "Longitude": [-112.1871, -83.5532, -119.5383, -110.5885]
                # make a dataframe and plot the USA map with the data as annotated dots
               df = pd.DataFrame(data)
                # define the desired colors
               colors = ["limegreen", "dodgerblue", "orange", "red"]
               # generate a scatter plot on a geographical map of the USA, with customized appearance and layout settings
fig = px.scatter_geo(df, lat="Latitude", lon="Longitude", hover_name="Location")
fig.update_geos(visible=True, resolution=110, scope="usa", showcountries=True, countrycolor="black", showsubunits=True, subunitcolor="gray", landcolor="#f8f8f8")
fig.update_layout(autosize=True, margin={"r":0,"t":20,"l":0,"b":0})
               # customize the dot colors
fig.update_traces(marker=dict(size=8, color=colors), selector=dict(type="scattergeo"))
                # Location annotation
                fig.add_trace(
                      go.Scattergeo(
                             lat=df["Latitude"] + 1,
lon=df["Longitude"],
                            ""cb>Bryce Canyon, Utah",
  "cb>Bryce Canyon, Utah</p",
  "cb>Great Smoky Mountains, Tennessee",
  "cb>Yosemite, California",
  "cb>Yosemite, California",
  "cb>Yosemite, California",
  "cb>Yosemite, California",
  "cb>Yosemite, California",
  "cb>Yosemite, California
                                    "<b>Yellowstone, Wyoming</b>"
                             mode="text",
textposition="middle center",
                             showlegend=False,
hoverinfo="none",
                              textfont=dict(size=12, color=colors),
```

```
# title
fig.add_annotation(
    x=0.5,
    y=1.05,
    text="\dots\b\b\biodiversity\ in Focus: Exploring Species Endangerment and Sightings across Four American National Parks</b>",
    showarrow=False,
    font=dict(size=16, color="black", family="Calibri"),
)

# lock the layout
fig.update_layout(dragmode=False, mapbox=dict(zoom=False))

# save the figure
fig.write_image("TitleFigure1.png", scale=3)

# display the figure
fig.show()
```

Biodiversity in Focus: Exploring Species Endangerment and Sightings across Four American National Parks



National Park Surface Areas

Yellowstone National Park

Approximately 8991 km² (3471 square miles).

Yosemite National Park

Approximately 3027 km² (1168 square miles).

Great Smoky Mountains National Park

Approximately 2113 km² (815 square miles).

Bryce Canyon National Park

Approximately 145 km² (56 square miles).

Title Figure 2: National Park Surface Area Comparison

```
In [53]: # adjust the plot size
plt.subplots(figslze(8, 6))

# park names
parks = ["Yellowstone", "Yosemite", "Great Smoky Mountains", "Bryce Camyon"]

# park names
areas_km = [8991, 3027, 2113, 145]

# set colors for the bubbles
colors = ["red", "orange", "dodgerblue", "limegreen"]

# create the bubble chart with switched axes
plt.scatter([1] * In(nparks), parks, s=areas_km, excolors, edgecolor="black")

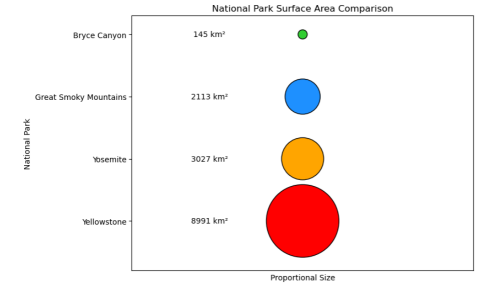
# add Labets and title
plt.valabel("Proportional Size")
plt.viabel("Proportional Park")
plt.title("Mational Park" Surface Area Comparison")

# set the y-axis Limit and tick spacing
plt.vjin(=0.8, 3.3)
plt.yticks(range(len(parks)), parks)

# remove x-axis ticks
plt.xticks([])

# annotate each scatter point with the corresponding areas_km value
for i, area in enumerate[areas_km]:
plt.text(1-0.8), i, "f'area jkm", han"center", va="center")

# sove the fjoure
plt.show()
```



Yellowstone National Park is a breathtaking natural wonder located in Wyoming

Renowned for its geothermal features, including the iconic Old Faithful geyser. Yellowstone is the first national park in the United States and is recognized for its diverse wildlife and stunning landscapes.

Yosemite National Park is a captivating destination in California.

Home to towering granite cliffs, cascading waterfalls, and ancient sequoia groves. Yosemite is celebrated for its awe-inspiring beauty and outdoor recreation opportunities.

Great Smoky Mountains National Park is a cherished treasure nestled between Tennessee and North Carolina.

This park is renowned for its mist-covered mountains, rich biodiversity, and vibrant fall foliage, making it one of the most visited national parks in the USA.

Bryce Canyon National Park is an extraordinary geological marvel situated in Utah.

Characterized by its distinctive orange-hued rock formations called hoodoos, Bryce Canyon offers unparalleled panoramic vistas and an opportunity to explore the unique beauty of the high desert landscape.

5: Questions and Analysis

5.1: Which species were spotted the most at each park?

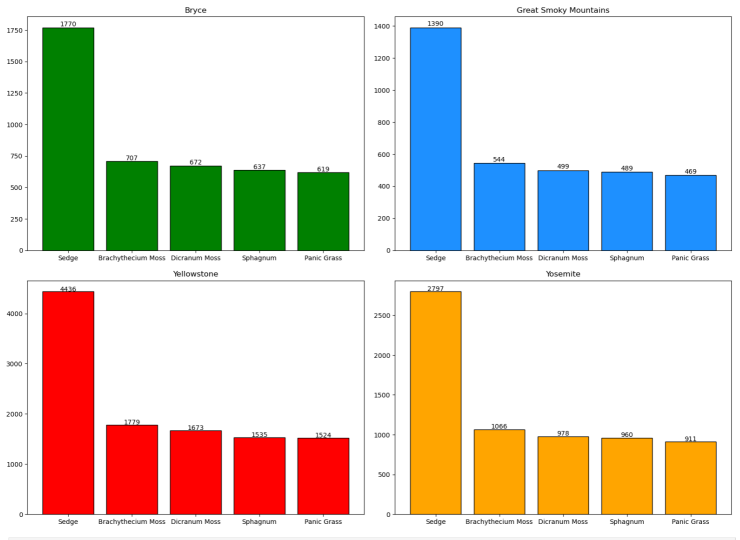
```
In [54]: # most observed species per category
          species_per_park = biodiversity.groupby(["park_name", "category", "common_names"]).observations.sum()
          # display the top 5 observed species in each park
for park in species_per_park.index.levels[0]:
              sorted_per_park = species_per_park[park].sort_values(ascending=False)
              print(f"National Park: {park}")
              print(sorted_per_park.head(5), "\n")
        National Park: Bryce
        category
Vascular Plant
                              common_names
                              Sedge
        Nonvascular Plant Dicranum Moss
                                                       707
                                                       672
                              Brachythecium Moss
                              Bryum Moss
                                                        637
        Name: observations, dtype: int64
        National Park: Great Smoky Mountains
                          common_names
        category
Vascular Plant
        Vascular Plant Sedge
Nonvascular Plant Dicranum Moss
                                                       1390
                             Brachythecium Moss
                                                       499
        Vascular Plant
                             Panic Grass
        Nonvascular Plant Bryum Moss
Name: observations, dtype: int64
                                                       469
        National Park: Yellowstone
        category
Vascular Plant
                              common_names
Sedge
        Nonvascular Plant Dicranum Moss
                                                      1779
                              Brachythecium Moss
                             Sphagnum
Panic Grass
                                                       1535
        Vascular Plant
                                                      1524
        Name: observations, dtype: int64
        National Park: Yosemite
        category
Vascular Plant
                              common_names
                              Sedge
                                                       2797
        Nonvascular Plant Brachythecium Moss
                                                       1066
                             Dicranum Moss
                                                       978
                              Sphagnum
                                                        960
        Vascular Plant
                              Panic Grass
                                                       911
        Name: observations, dtype: int64
```

Figure 1: Most Observed Species per National Park

```
In [55]: # this observations distribution is another indicator that this dataset is fictional!
            # define the parks and subplots layout
           parks = species_per_park.index.levels[0]
num_parks = len(parks)
           # define colors for each park
colors = ["green", "dodgerblue", "red", "orange"]
           # create the figure and subplots
fig, axes = plt.subplots(2, 2, figsize=(16, 12))
axes = axes.flatten()
           # iterate over each park
for i, park in enumerate(parks):
                sorted_species = species_per_park.loc[park].sort_values(ascending=False)
                # get the top 5 species for the current park
top_species = sorted_species.head(5)
                 # generate the bar graph
                ax = axes[i]
                ax.bar(top_species.index.get_level_values("common_names"), top_species.values,
                         edgecolor="black", color=colors[i])
                # add annotations on top of each bar
                for j, v in enumerate(top_species.values):
                     ax.text(j, v + 10, str(v), ha="center")
                ax.set title(park)
           # set the title for the whole figure
fig.suptitle("Most Observed Species per National Park", fontsize=16)
              set x-axis tick positions and labels
            for ax in axes:
                ax.set_xticks(range(len(top_species.index)))
                ax.set_xticklabels(top_species.index.get_level_values("common_names"))
            # hide empty subplots
           if num_parks < 4:
    for i in range(num_parks, 4):
        fig.delaxes(axes[i])</pre>
            # adjust the spacing between subplots
           fig.tight_layout(pad=1.5)
            # save the figure
           plt.savefig("Figure1.png", dpi=300, bbox_inches="tight")
            # display the figure
```

plt.show()

Most Observed Species per National Park



```
American Bullfrog
Pickerel Frog
                                   677
        Marbled Salamander
        Eastern Mud Salamander
                                   656
        Mud Salamander
        Name: observations, dtype: int64
        Category: Bird
        common_names
Eurasian Collared-Dove 1785
        Water Pipit
                                  1728
        Brewster's Warbler
                                  1704
        Chestnut-Sided Warbler
                                  1310
        Name: observations, dtype: int64
        Category: Fish
        common names
        Brook Trout
                            1270
        Spotfin Shiner
                            1140
        Mottled Sculpin
        Whitetail Shiner
                            1120
        Blacktail Shiner
                            1119
        Name: observations, dtype: int64
        Category: Mammal
        Uinta Chipmunk
                           1850
        Panther
                           1711
        Common Raccoon
        Mink
                           1644
        Name: observations, dtype: int64
        Category: Nonvascular Plant
        common_names
        Dicranum Moss
        Brachythecium Moss 3910
        Bryum Moss
        Sphagnum
                             3476
                              2955
        Name: observations, dtype: int64
        Category: Reptile
        common_names
        Sierra
                                  1257
        Rubber Boa
        Western Painted Turtle
                                   669
        California Nightsnake
        Corn Snake
                                   649
        Name: observations, dtype: int64
        Category: Vascular Plant
        common_names
        Sedge
                            3543
        Panic Grass
        Bladder Campion
                             2886
        Goldenrod
                             2860
        Goosefoot Violet
                             2784
        Name: observations, dtype: int64
         Figure 2: Top Five Most Observed Species per Category
In [57]: # define the categories and subplots Layout
         categories = species_per_category.index.levels[0]
         num_categories = len(categories)
         num_plots_per_col = (num_categories + num_plots_per_row - 1) // num_plots_per_row
         # create the figure and subplots
fig, axes = plt.subplots(num_plots_per_col, num_plots_per_row, figsize=(18, 12))
axes = axes.flatten()
```

print(f"Category: {category}")
print(sorted_per_category.head(5), "\n")

Category: Amphibian common names

```
# iterate over each category
for i, category in enumerate(categories):
    sorted_species = species_per_category[category].sort_values(ascending=False)
    # get the top 5 species for the current category
    top_species = sorted_species.head(5)
    # generate the bar graph
    ax = axes[i]
    ax.bar(top_species.index, top_species.values, color="dodgerblue", edgecolor="black")
    # add annotations on top of each bar
    for j, v in enumerate(top_species.values):
        ax.text(j, v + 10, str(v), ha="center")
    ax.set_title(category)
    # set the title for the whole figure
fig.suptitle("Most Observed Species per Category", fontsize=16)
    # set x-axis tick positions and Labels
    ax.set_xticks(range(len(top_species.index)))
    ax.set_xticklabels(top_species.index, rotation=30)
 # hide empty subplots
if num_categories < num_plots_per_row * num_plots_per_col:</pre>
    for i in range(num_categories, num_plots_per_row * num_plots_per_col):
        fig.delaxes(axes[i])
fig.tight_layout(pad=1)
plt.savefig("Figure2.png", dpi=300, bbox_inches="tight")
```



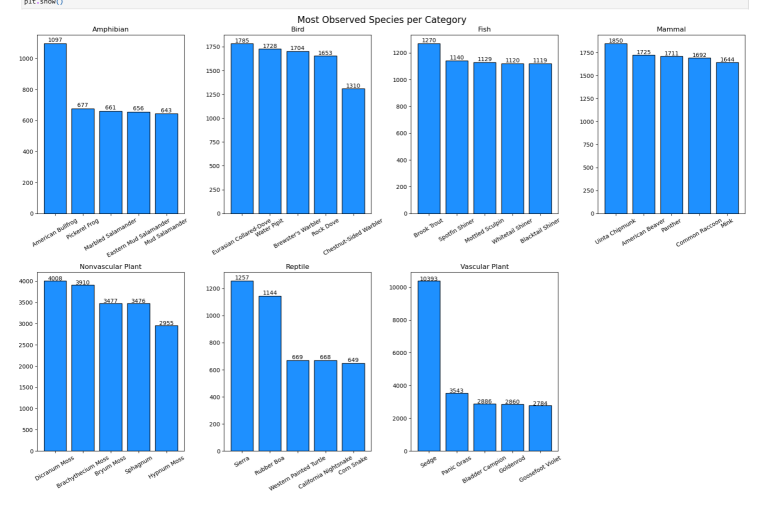


Figure 3: Total Observations per Category

```
In [58]: # adjust the plot size
plt.subplots(figsize=(8, 6))
          # sum and sort the observations for each category
total_observations_per_category = biodiversity.groupby("category")["observations"].sum().reset_index().sort_values("observations", ascending=False)
           # divide by thousand for readability
total_observations_per_category["observations"] = total_observations_per_category["observations"] / 1000
           # determine the color map
           cmap = sns.color_palette("coolwarm_r", len(total_observations_per_category))
           # barplot
           sns.barplot(data=total\_observations\_per\_category, \ x="category", \ y="observations", \ ec="k", \ palette=cmap)
           plt.title("Total Observations per Category")
plt.xlabel("Category")
           plt.ylabel("Number of Observations (thousands)")
           # add annotation for readability
           for index, value in enumerate(total_observations_per_category["observations"]):
                plt.text(index, value, f"{(value/total_observations_per_category.observations.sum() * 100):.0f}%", ha="center", va="bottom")
           # format x and y-axis Labels format = ticker.FuncFormatter(lambda x, pos: f''(x:.0f)K'')
           plt.gca().yaxis.set_major_formatter(format)
plt.xticks(rotation=25)
           plt.savefig("Figure3.png", dpi=300, bbox_inches="tight")
           # display the figure
           plt.show()
```

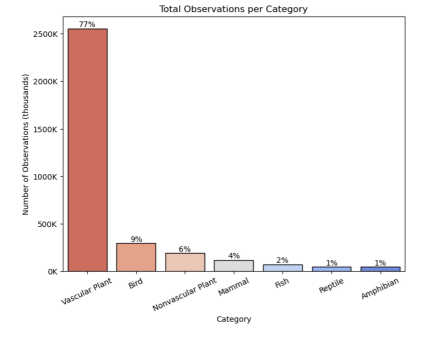


Figure 4: Total Observations per National Park

```
In [59]: # adjust the plot size
plt.subplots(figsize=(8, 6))
                              # sum and sort the observations for each park
                            total\_observations\_per\_park = biodiversity.groupby ("park\_name") ["observations"].sum().reset\_index().sort\_values ("observations", ascending=False) ["observations"].sum().sort\_values ("observations").sum().sort\_values ("observations", ascending=False) ["observations"].sum().sort\_values ("observations", ascending=False) ["observations"].sum().sort\_values ("observations").sum().sort\_values ("observations").sum().sort\_values ("observations").sum().sort\_
                            # divide by thousand for readability
total_observations_per_park["observations"] = total_observations_per_park["observations"] / 1000
                            # set custom colors for each bar
colors = ["red", "orange", "dodgerblue", "limegreen"]
                             # barplot with custom colors
                             sns.barplot(data=total_observations_per_park, x="park_name", y="observations", ec="k", palette=colors)
                            plt.title("Total Observations per National Park")
plt.xlabel("National Park")
                            plt.ylabel("Number of Observations (thousands)")
                            # add annotation for easier graph reading
for index, value in enumerate(total_observations_per_park["observations"]):
                                        # format y-axis labels format = ticker.FuncFormatter(lambda x, pos: f''(x:.0f)K'')
                            plt.gca().yaxis.set_major_formatter(format)
                           # save the figure
plt.savefig("Figure4.png", dpi=300, bbox_inches="tight")
                              # display the figure
                            plt.show()
```

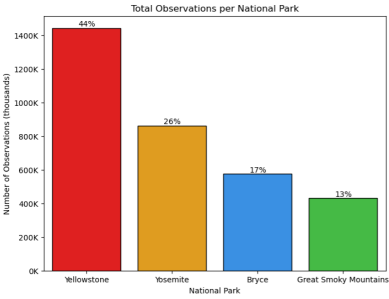


Figure 5: Observations per Category per National Park

```
pivot the dataframe to have park names as rows, category as columns, and observations as values
 pivoted_data = grouped_data.pivot(index="park_name", columns="category", values="observations")
 # specify the desired order of parks for sorting
park_order = ["Yellowstone", "Yosemite", "Bryce", "Great Smoky Mountains"]
   sort the pivoted data DataFrame based on the specified park order
 pivoted_data = pivoted_data.loc[park_order]
       rt the columns within each park in descending orde
 pivoted_data = pivoted_data[pivoted_data.sum().sort_values(ascending=False).index]
  # create a figure with two subplots
 fig, axes = plt.subplots(1, 2, figsize=(15, 6))
 # plot the first subplot
 ax1 = axes[0]
 pivoted_data.plot(kind="bar", stacked=False, edgecolor="black", width=0.8, ax=ax1)
 # format x and y-axis
 ax1.yaxis.set_major_formatter(ticker.FuncFormatter(lambda x, pos: "{:.0f}K".format(x * 1e-3)))
 ax1.set_xticklabels(pivoted_data.index, rotation=0)
 # add axis labels, title, and legend
ax1.set_xlabel("National Park")
 ax1.set_ylabel("Number of Observations (thousands)")
ax1.set_title("Observations per Category per National Park")
ax1.legend(title="Category", frameon=False, fontsize=8)
 # plot the second subplot with logarithmic y-scale
 ax2 = axes[1]
 pivoted_data.plot(kind="bar", stacked=False, edgecolor="black", width=0.8, ax=ax2)
 # format x and y-axis with Logarithmic scale
ax2.yaxis.set_major_formatter(ticker.FuncFormatter(lambda x, pos: "{:.0f}K".format(x * 1e-3)))
 ax2.set_xticklabels(pivoted_data.index, rotation=0)
# logarithmic scale allows for a better understanding of the relationship between the categories in each park.
 # It is important to address that this distorts the bigger picture.
# For example, Yellowstone has the majority of observations across all categories, especially for vascular plants.
 ax2.set_yscale("log")
 # add axis Labels, title, and Legend
ax2.set_xlabel("National Park")
ax2.set_ylabel("Number of Observations")
ax2.set_title("Observations per Category per National Park (Logarithmic Scale)")
 ax2.legend(title="Category", frameon=False, fontsize=8, loc=(0.78,0.706))
 # adjust the spacing between subplots
 plt.tight layout(pad=2.5)
 plt.savefig("Figure5.png", dpi=300, bbox_inches="tight")
 # display the figure
 plt.show()
                                                                                                                                       Observations per Category per National Park (Logarithmic Scale)
                                Observations per Category per National Park
                                                                                                Category
                                                                                                                                                                                                                       Category
                                                                                                                           106
                                                                                                                                                                                                                      Vascular Pla
Bird
Nonvascular
   1000K
                                                                                               Mammal
                                                                                                                                                                                                                      Mammal
                                                                                                                                                                                                                 Mam
Fish
                                                                                               Fish
                                                                                               Reptile
                                                                                                                                                                                                                  Reptile
Number of Observations (thousands)
                                                                                                                                                                                                                  Amphibian
    800K
                                                                                                                        of Observations
                                                                                                                           105
    600K
                                                                                                                        Number
     400K
    200K
                                                                                                                           10
       0K
                                                                                    Great Smoky Mountains
                                                                                                                                          Yellowstone
                     Yellowstone
                                              Yosemite
                                                                       Bryce
                                                                                                                                                                   Yosemite
                                                                                                                                                                                            Bryce
                                                                                                                                                                                                         Great Smoky Mountains
```

Figure 6: Species Distribution by Category

National Park

```
In [61]: # adjust the plot size
    plt.subplots(figsize(8, 6))

# count species by category
    species_category = pd.DataFrame(biodiversity.category.value_counts())

# calculate the percentage of each category
    species_category["percentage"] = (species_category["count"] / species_category["count"].sum() * 100)

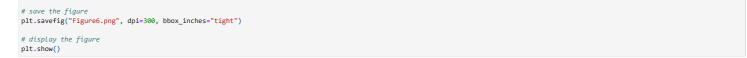
# x axis is for categories
    x = species_category.index

# determine the color map
    cmap = sns.light_palette("blue", len(species_category), reverse=True)

# seaborn bar plot
    sns.barplot(data=species_category, x=x, y="percentage", ec="k", palette=cmap)
    plt.title("Species Distribution by Category")
    plt.ylabel("Category")
    plt.ylabel("Percent / %")
    plt.xticks(rotation=25)

# add annotation for easier graph reading
    for index, value in enumerate(species_category["percentage"]):
        plt.text(index, value, f"(value:.2f)%", has "center", va="bottom")
```

National Park



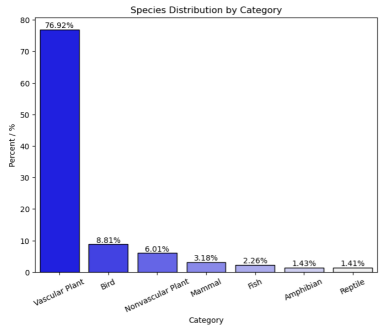


Table 1: Species Distribution by Category: Count and Percentage

```
In [62]: display(species_category.style.set_caption("Species Distribution by Category: Count and Percentage"))
       Species Distribution by Category: Count and
                   Percentage
                        count percentage
               category
           Vascular Plant 17048 76 917524
                   Bird 1952
       Nonvascular Plant 1332
                                6.009746
               Mammal 704 3.176322
                   Fish 500
                                 2.255910
             Amphibian
                         316
                                 1.425735
                 Reptile 312
                                 1.407688
```

5.2: What is the Distribution of Conservation Status for Animals?

Figure 7: Overall Conservation Status Distribution (Logarithmic scale)

```
In [63]: # adjust the plot size
           plt.subplots(figsize=(8, 6))
           conservation_status = pd.DataFrame(biodiversity.conservation_status.value_counts())
            # calculate and display the percentage of each category
           conservation_status["percentage"] = (conservation_status["count"] / conservation_status["count"].sum() * 100)
display(conservation_status.style.set_caption("Overall Conservation Status Distribution: Count and Percentage"))
            # x axis is for conservation status
            x = biodiversity.conservation_status.unique()
            # determine the color map
           cmap = sns.light_palette("magenta", len(conservation_status))
            # seaborn bar plot
            \verb|sns.barplot(data=conservation_status, x=x, y="percentage", ec="k", palette=cmap)|
           plt.title("Overall Conservation Status Distribution (Logarithmic scale)")
plt.xlabel("Conservation Status")
           plt.ylabel("Percent / %")
            # add annotation for easier graph reading
           for index, value in enumerate(conservation_status["percentage"]):
    plt.text(index, value, f"{value:.2f}%", ha="center", va="bottom")
            # logarithmic scale for better readability
           plt.yscale("log")
           # save the figure
plt.savefig("Figure7.png", dpi=300, bbox_inches="tight")
```

```
display the figure
 plt.show()
 Overall Conservation Status Distribution:
        Count and Percentage
                  count percentage
conservation status
     Least concern 21452 96.787583
Species of Concern 604
                          2.725140
      Endangered
                           0.270709
      Threatened 36 0.162426
      In Recovery
                    12
                           0.054142
                 Overall Conservation Status Distribution (Logarithmic scale)
              96.79%
    10<sup>2</sup>
```

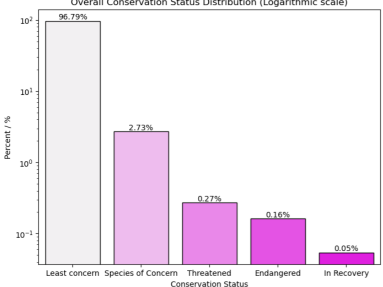


Table 2: Conservation Status Distribution per Park

```
In [64]: # counting conservation status p
          conservation_status_per_park = pd.DataFrame(biodiversity.groupby(["park_name", "conservation_status"]).size().unstack(fill_value=0))
          # there is no need to plot this beacuse the amount of each conservation status is exactly the same for each national park;
          # This is a fine indication that this is a fictional dataset!
display(conservation_status_per_park.style.set_caption("Conservation Status Distribution per Park"))
                                       Conservation Status Distribution per Park
             conservation_status Endangered In Recovery Least concern Species of Concern Threatened
                          Brvce
                                                                     5363
                                                                                           151
                                                                                                          9
         Great Smoky Mountains
                                           15
                                                                     5363
                                                                                           151
                                                                                                          9
                                           15
                                                         3
                                                                     5363
                                                                                           151
                                                                                                          9
                    Yellowstone
                                           15
                                                                     5363
                                                                                           151
```

Figure 8: Conservation Status Distribution per Category (Logarithmic scale)

```
In [65]: #
            create subplots with desired figure size
          fig, ax = plt.subplots(figsize=(8, 6))
          # count the conservation status for each catego
          conservation_status_per_category = biodiversity.groupby(["category", "conservation_status"]).size().unstack(fill_value=0)
          # sort each category by conservation status occurrence from highest to lowest
          conservation\_status\_per\_category = conservation\_status\_per\_category.reindex(conservation\_status\_per\_category.sum(axis=1).sort\_values(ascending=False).index)
          # sort each conservation status within each category
          conservation_status_per_category = conservation_status_per_category[conservation_status_per_category.sum().sort_values(ascending=False).index]
          conservation_status_per_category.plot(kind="barh", stacked=False, edgecolor="black", width=0.8, ax=ax)
          # customizing the plot
ax.set_title("Conservation Status Distribution per Category (Logarithmic scale)")
          ax.set_xlabel("Count")
ax.set_ylabel("Category
          # Logarithmic scale for better readability
          ax.set_xscale("log")
          ax.legend(frameon=False)
# adjusting the height to increase the space between each bar group
          ax.set_ylim([-0.55, len(conservation_status_per_category)-0.9])
          ax.invert_yaxis()
          plt.legend(frameon=False, title="Conservation Status")
          plt.savefig("Figure8.png", dpi=300, bbox_inches="tight")
          # display the figure
          plt.show()
```

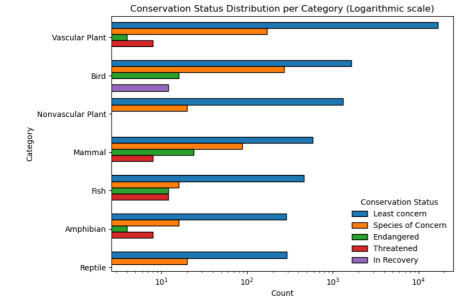
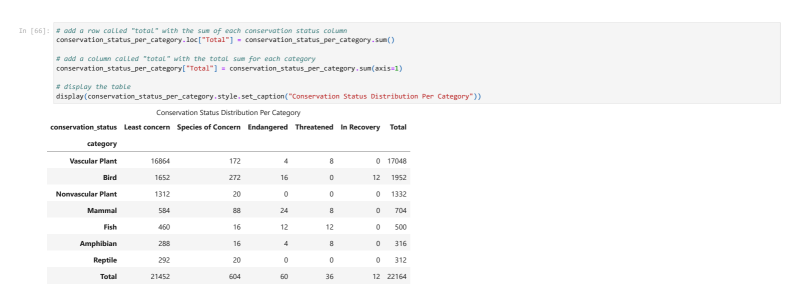


Table 3: Conservation Status Distribution Per Category



5.3: Are certain types of species more likely to be endangered?

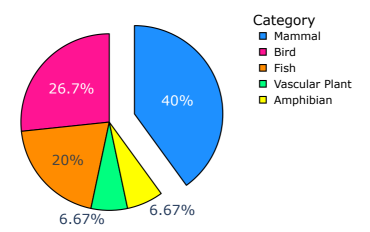
Table 4: Conservation Status Distribution Per Category in Percentages

```
In [67]: # table to calculate the percentage of conservation status per category
        endangerment_table = conservation_status_per_category.copy().drop(labels="Total", axis=1)
        # calculate the percentage for each row
endangerment_table_percent = endangerment_table.copy()
        for column in endangerment_table.columns:
                display(endangerment_table_percent.style.set_caption("Conservation Status Distribution per Category in Percentages").format("{:.2f}"))
                       Conservation Status Distribution per Category in Percentages
       conservation_status Least concern Species of Concern Endangered Threatened In Recovery
                category
            Vascular Plant
                                                              6.67
                                                                        22.22
                                                                                    0.00
                    Bird
                                 7.70
                                                 45.03
                                                             26.67
                                                                         0.00
                                                                                   100.00
        Nonvascular Plant
                                 6.12
                                                  3.31
                                                              0.00
                                                                         0.00
                                                                                     0.00
                Mammal
                                 2.72
                                                 14.57
                                                             40.00
                                                                        22.22
                                                                                    0.00
                    Fish
                                 2.14
                                                  2.65
                                                             20.00
                                                                        33.33
                                                                                    0.00
              Amphibian
                                 1.34
                                                  2.65
                                                              6.67
                                                                        22.22
                                                                                    0.00
                 Reptile
                                 1.36
                                                  3.31
                                                                         0.00
                                                                                    0.00
                                                              0.00
                                100.00
                                                 100.00
                                                            100.00
                                                                        100.00
                                                                                   100.00
```

Figure 9: Endangered Species Distribution

```
In [68]: # dataframe containing only endangered species
           endangered_species = biodiversity.loc[biodiversity["conservation_status"] == "Endangered"]
           # endangered species count per category
endangered_category_distribution = endangered_species.category.value_counts()
           colors = ["DodgerBlue", "DeepPink", "DarkOrange", "SpringGreen", "Yellow"]
           # pie chart with modifications
fig = go.Figure(data=[go.Pie(
                labels=endangered_category_distribution.index, values=endangered_category_distribution,
                pull=[0.3],
                insidetextorientation="horizontal")]
            # pie chart design
            fig.update_layout(width=800, height=600, title={"text": "Endangered Species Distribution", "font": {"size": 30, "color": "black"}})
           fig.update_traces(hoverinfo="label+value"
                                textinfo="percent",
                                 textfont size=25.
                                 marker=dict(colors=colors, line=dict(color="#000000", width=2))
            # nie chart desiar
           fig.update_layout(
                "text": "Endangered Species Distribution",
    "font": {"size": 30}},
legend=dict(
                     title="Category",
title_font=dict(size=25, color="black"),
font=dict(size=20, color="black"),
            # save the figure
           fig.write_image("Figure9.png", scale=2)
            # display the figure
           fig.show()
```

Endangered Species Distribution



```
In [69]: display(endangered_category_distribution)

category

Mammal 24

Bird 16

Fish 12

Vascular Plant 4

Amphibian 4

Name: count, dtype: int64

In [70]: endangered_mammals = biodiversity[(biodiversity["conservation_status"] == "Endangered") & (biodiversity["category"] == "Mammal")].sort_values("observations", ascending=False)

In [71]: display(endangered_mammals.head(10).style.set_caption("Top 10 Endangered Mammal Species"))
```

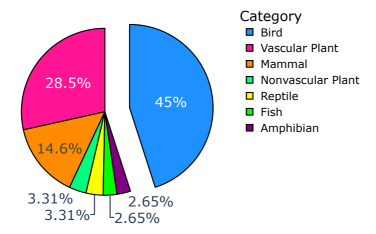
Top 10 Endangered Mammal Species

		•					
	park_name	scientific_name	category	common_names	$conservation_status$	observations	
11886	Yellowstone	Canis lupus	Mammal	Gray Wolf	Endangered	330	
17427	Yosemite	Canis lupus	Mammal	Gray Wolf	Endangered	196	
804	Bryce	Canis lupus	Mammal	Gray Wolf	Endangered	130	
14406	Yellowstone	Myotis sodalis	Mammal	Indiana Bat	Endangered	68	
14401	Yellowstone	Myotis grisescens	Mammal	Gray Myotis	Endangered	68	
13374	Yellowstone	Glaucomys sabrinus coloratus	Mammal	Carolina Northern Flying Squirrel	Endangered	67	
14574	Yellowstone	Ovis canadensis sierrae	Mammal	Sierra Nevada Bighorn Sheep	Endangered	67	
11887	Yellowstone	Canis rufus	Mammal	Red Wolf	Endangered	60	
6345	Great Smoky Mountains	Canis lupus	Mammal	Gray Wolf	Endangered	59	
19942	Yosemite	Myotis grisescens	Mammal	Gray Myotis	Endangered	39	

Figure 10: Species of Concern Distribution

```
In [72]: # dataframe containing only species of concern
           species_of_concern = biodiversity.loc[biodiversity["conservation_status"] == "Species of Concern"]
           # endangered species count per category
          species_of_concern_distribution_by_category = species_of_concern.category.value_counts()
          colors = ["DodgerBlue", "DeepPink", "DarkOrange", "SpringGreen", "Yellow", "Lime", "Purple"]
           # pie chart with modifications
           fig = go.Figure(data=[go.Pie(
               labels=species_of_concern_distribution_by_category.index, values=species_of_concern_distribution_by_category,
               pull=[0.3].
               insidetextorientation="horizontal")]
          fig.update_layout(width=800, height=600, title={"text": "Endangered Species Distribution", "font": {"size": 30, "color": "black"}})
fig.update_traces(hoverinfo="label+value",
                               textinfo="percent",
                               textfont size=25.
                               marker=dict(colors=colors, line=dict(color="#000000", width=2))
           # nie chart desiar
           fig.update_layout(
               title={
                   "test": "Species of Concern Distribution",
"font": {"size": 30}},
               legend=dict(
                   title="Category"
                    title_font=dict(size=25, color="black"),
font=dict(size=20, color="black"),
           # save the figure
           fig.write_image("Figure10.png", scale=2)
          fig.show()
```

Species of Concern Distribution



	Top 10 Species of Concern Bird Species						
	park_name	scientific_name	category	common_names	conservation_status	observations	
14608	Yellowstone	Pandion haliaetus	Bird	Osprey	Species of Concern	466	
14487	Yellowstone	Nycticorax nycticorax	Bird	Black-Crowned Night-Heron	Species of Concern	439	
15456	Yellowstone	Riparia riparia	Bird	Bank Swallow	Species of Concern	425	
13307	Yellowstone	Gavia immer	Bird	Common Loon	Species of Concern	408	
18848	Yosemite	Gavia immer	Bird	Common Loon	Species of Concern	319	
20149	Yosemite	Pandion haliaetus	Bird	Osprey	Species of Concern	282	
20997	Yosemite	Riparia riparia	Bird	Bank Swallow	Species of Concern	270	
20028	Yosemite	Nycticorax nycticorax	Bird	Black-Crowned Night-Heron	Species of Concern	269	
13953	Yellowstone	Limosa fedoa	Bird	Marbled Godwit	Species of Concern	266	

Figure 11: Threatened Species Distribution

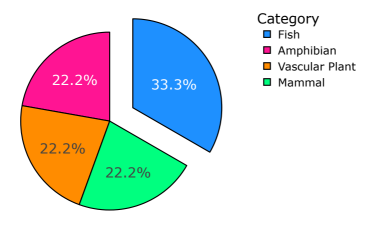
Bird

13192 Yellowstone Falco columbarius

```
In [76]: # dataframe containing only species of concern
          threatened_species = biodiversity.loc[biodiversity["conservation_status"] == "Threatened"]
          # endangered species count per category
threatened_species_by_category = threatened_species.category.value_counts()
          colors = ["DodgerBlue", "DeepPink", "DarkOrange", "SpringGreen", "Yellow", "Lime", "Purple"]
          # pie chart with modifications
fig = go.Figure(data=[go.Pie(
               labels=threatened_species_by_category.index,
values=threatened_species_by_category,
               pull=[0.3],
               insidetextorientation="horizontal")]
           # nie chart desian
           fig.update_layout(width=800, height=600, title={"text": "Endangered Species Distribution", "font": {"size": 30, "color": "black"}})
          textfont size=25.
                               marker=dict(colors=colors, line=dict(color="#000000", width=2))
           # pie chart design
           fig.update_layout(
               .update_layout(
title={
    "text": "Threatened Species Distribution",
    "font": {"size": 30}},
legend=dict(
                    title="Category",
title_font=dict(size=25, color="black"),
                    font=dict(size=20, color="black"),
          fig.write_image("Figure11.png", scale=2)
           # display the figure
          fig.show()
```

Merlin Species of Concern

Threatened Species Distribution



	Top 10 Threatened Fish Species						
	park_name	scientific_name	category	common_names	conservation_status	observations	
14477	Yellowstone	Noturus flavipinnis	Fish	Yellowfin Madtom	Threatened	126	
14510	Yellowstone	Oncorhynchus clarkii henshawi	Fish	Lahontan Cutthroat Trout	Threatened	111	
13043	Yellowstone	Erimonax monachus	Fish	Spotfin Chub	Threatened	109	
20051	Yosemite	Oncorhynchus clarkii henshawi	Fish	Lahontan Cutthroat Trout	Threatened	85	
20018	Yosemite	Noturus flavipinnis	Fish	Yellowfin Madtom	Threatened	72	
18584	Yosemite	Erimonax monachus	Fish	Spotfin Chub	Threatened	69	
3428	Bryce	Oncorhynchus clarkii henshawi	Fish	Lahontan Cutthroat Trout	Threatened	57	
3395	Bryce	Noturus flavipinnis	Fish	Yellowfin Madtom	Threatened	55	
8969	Great Smoky Mountains	Oncorhynchus clarkii henshawi	Fish	Lahontan Cutthroat Trout	Threatened	48	
1961	Bryce	Erimonax monachus	Fish	Spotfin Chub	Threatened	46	

5.4: Are the differences between species and their conservation status significant?

Figure 12: Species Observations per Conservation Status

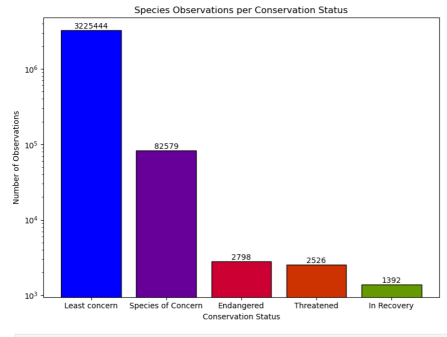
```
In [80]: # sum the number of observations for each conservation status
observations_per_conservation_status = biodiversity.groupby("conservation_status")["observations"].sum().sort_values(ascending=False)

# color mapped bar chart
plt.figure(figsize=(8, 6))
colors = plt.cm.brg_r(np.linspace(1, 0.2, len(observations_per_conservation_status)))
plt.bar(observations_per_conservation_status.index, observations_per_conservation_status.values, edgecolor="black", color=colors)
plt.valabel("Conservation Status")
plt.valabel("Number of Observations")
plt.title("Species Observations per Conservation Status")
plt.tight_layout()
plt.yscale("log")

# add annotations for readability
for index, value in enumerate(observations_per_conservation_status):
    plt.title(index, value, f"{value:.0f}", ha="center", va="bottom")

# save the figure
plt.savefig("Figure12.png", dpi=300, bbox_inches="tight")

# display the figure
plt.show()
```



```
In [81]: observations_per_conservation_status

Out[81]: conservation_status
    Least concern 3225444
    Species of Concern 82579
    Endangered 2798
    Threatened 2526
    In Recovery 1392
    Name: observations, dtype: int64
```

Figure 13: Conservation Status Distribution per Category

```
# sort each category by conservation status occurrence from highest to Lowest
conservation_status_per_category = conservation_status_per_category.reindex(conservation_status_per_category.sum(axis=1).sort_values(ascending=False).index)

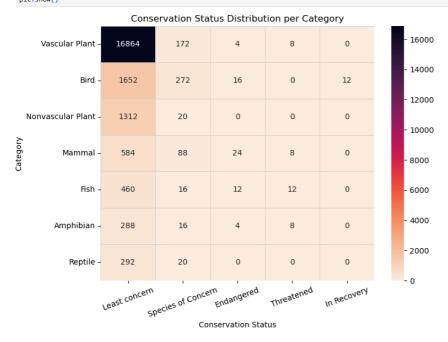
# sort each conservation status within each category
conservation_status_per_category = conservation_status_per_category[conservation_status_per_category.sum().sort_values(ascending=False).index]

# create a heatmap
fig, ax = plt.subplots(figsize=(8, 6))
sns.heatmap(conservation_status_per_category, annot=True, fmt="d", cmap="rocket_r", cbar=True, ax=ax, linewidths=0.5, linecolor="lightgray")

# customizing the plot
ax.set_title("Conservation Status Distribution per Category")
ax.set_title("Conservation Status")
ax.set_title("Conservation Status")
ax.set_title(ax.get_xticklabels(), rotation=20)

# save the figure
plt.savefig("Figure13.png", dpi=300, bbox_inches="tight")

# display the figure
plt.show()
```



5.4.1: Shannon-Weaver and Simpson's Index for each National Park

```
# calculate the Shannon-Weaver index for each National Park
shannon_indices = biodiversity.groupby("park_name")["observations"].apply(lambda x: entropy(x.value_counts(normalize=True)))

# calculate the Simpson's index for each National Park
simpsons_indices = biodiversity.groupby("park_name")["observations"].apply(lambda x: 1 - np.sum((x.value_counts() / len(x)) ** 2))
```

Table 5: Shannon-Weaver and Simpson's Index for each National Park

```
In [84]: # print the calculated indices
         print(round(shannon_indices, 4))
         print("\nSimpson's Index:")
         print(round(simpsons_indices, 4))
        Shannon-Weaver Index:
        park name
                                 4.6192
        Bryce
        Great Smoky Mountains
                                 4.5724
        Yellowstone
        Yosemite
                                 4.6353
        Name: observations, dtype: float64
        Simpson's Index:
        park_name
                                 0.9873
        Bryce
        Great Smoky Mountains
                                 0.9871
        Yellowstone
                                 0.9875
        Yosemite
                                 0.9873
        Name: observations, dtype: float64
```

Both the Shannon-Weaver and Simpson's indices are used to measure biodiversity, but they provide slightly different perspectives on the diversity within each park.

The Shannon-Weaver index takes into account both species richness (the number of species present) and evenness (how evenly distributed the individuals are among species). A higher Shannon-Weaver index value indicates higher biodiversity. In this case, all four parks have relatively high Shannon-Weaver index values. This suggests that all the parks exhibit considerable species richness and evenness in terms of the observed species.

On the other hand, the Simpson's index focuses more on species dominance. It measures the probability that two individuals randomly selected from the population belong to the same species. The Simpson's index ranges from 0 to 1, with higher values indicating lower diversity and higher dominance of a few species. In this case, all four parks have high Simpson's index values, indicating a relatively low dominance of any specific species and a high level of diversity.

Overall, these results suggest that all four parks have relatively high biodiversity, with a diverse array of species present and a relatively even distribution of individuals among species.

5.4.2: Chi-squared test and Fisher's, Barnard's and Boschloo's exact tests

• Chi-squared (χ²) Test:

(https://en.wikipedia.org/wiki/Chi-squared_test)

The chi-square test is used to determine the association between two categorical variables. It compares the observed frequencies in a contingency table with the expected frequencies under the assumption of independence. The test statistic, χ^2 , is calculated based on the differences between observed and expected frequencies. The p-value is then derived from the chi-square distribution with appropriate degrees of freedom. A low p-value suggests that there is a significant association between the variables.

Fisher's Exact Test:

(https://en.wikipedia.org/wiki/Fisher%27s_exact_test)

Fisher's exact test is also used to analyze the association between categorical variables. It is commonly used when the sample size is small, or when any expected cell frequency in a contingency table is less than 5. Fisher's exact test calculates the probability of obtaining the observed distribution and all other possible distributions that are more extreme, assuming independence. The p-value represents the sum of these probabilities. A low p-value indicates a significant association between the variables.

Barnard's Exact Test:

(https://en.wikipedia.org/wiki/Barnard%27s_test)

Barnard's exact test is similar to Fisher's exact test and is used for analyzing the association between categorical variables. It calculates the probability of observing a given table, assuming independence, and compares it to the probability of observed table under a specific alternative hypothesis. The p-value is computed by summing the probabilities of tables that are at least as extreme as the observed table. Again, a low p-value indicates a significant association.

Boschloo's Exact Test:

(https://en.wikipedia.org/wiki/Boschloo%27s_test)

Boschloo's exact test is an alternative to Fisher's exact test that can be used when there are ordered categories or ordinal data. It tests for the association between two ordinal variables. The test calculates the probability of observing the data under the null hypothesis of no association, and the p-value is derived accordingly. A low p-value suggests a significant association between the ordinal variables.

Boschloo's exact test is not considered in this analysis due to its use for ordinal categories.

The operational difference between Barnard's exact test and Fisher's exact test is how they handle nuisance parameter(s) in calculating the p-value. Fisher's exact test avoids estimating nuisance parameters by conditioning on both margins, while Barnard's test considers all legitimate possible values and chooses the value that maximizes the p-value.

The theoretical difference is that Barnard's test uses the double-binomially distributed distribution, while Fisher's test uses the hypergeometric distribution for conditioning.

Both tests are valid, bounding the type I error rate at the alpha level. However, Barnard's test can be more powerful than Fisher's test by considering more extreme tables and not conditioning on the second margin, which Fisher's test ignores.

It is considered that Fisher's exact test is a more uniformly powerful alternative to the Chi-squared test, while Barnard's and Boschloo's exact tests are uniformly more powerful alternatives to Fischer's exact test

To analyze the statistics and p-values of these tests, usually the obtained p-value is compared to a significance level (such as 0.05; 5%). If the p-value is smaller than the significance level, the null hypothesis is rejected and it is concluded that there is evidence of an association between the variables. Conversely, if the p-value is larger than the significance level, the null hypothesis stands and it can be concluded that there is insufficient evidence to suggest an association.

 $In \ statistics, the \ \textbf{Bonferroni correction} \ is \ \textbf{a} \ \textbf{method to counteract the multiple comparisons problem} \ (https://en.wikipedia.org/wiki/Bonferroni_correction).$

Statistical hypothesis testing is based on rejecting the null hypothesis if the likelihood of the observed data under the null hypotheses is low.

If multiple hypotheses are tested, the probability of observing a rare event increases, and therefore, the likelihood of incorrectly rejecting a null hypothesis (i.e., making a Type I error) increases.

For these reasons, **p-value is corrected according to the Bonferroni correction**.

Number of 2x2 contingency tables that can be constructed without including same-named ones and inverse pairs (if we have A and B, B and A is not needed) involves selecting combinations of categories, we can calculate the number of unique combinations using the formula for combinations:

nCr = n! / (r!(n-r)!)

Where n is the total number of categories (7 in this case) and r is the number of categories to be selected (2 for a 2x2 table)

Applying this formula, we get

```
nCr = 7! / (2!(7-2)!) = 7! / (2!5!) = (7x6) / (2x1) = 21
```

Null Hypothesis: There are no significant differences between the protected and unprotected species

Alternative Hypothesis: There are significant differences between the protected and unprotected species.

To analyze the given results, we need to compare the p-values with the predetermined significance level of 0.00238. The significance level, also known as alpha (a), represents the threshold below which we reject the null hypothesis.

```
In [86]: protected_data = pd.DataFrame(protected_count).T
protected_data["protected_percentage"] = protected_data["protected"] / protected_data["total"] * 100
protected_data = protected_data.sort_values("protected_percentage", ascending=False)
display(protected_data.style.set_caption("Category Protection Breakdown: Counts, Totals, and Percentages"))

Category Protection Breakdown: Counts, Totals, and Percentages
```

	$not_protected$	protected	total	protected_percentage
Mammal	584	120	704	17.045455
Bird	1652	300	1952	15.368852
Amphibian	288	28	316	8.860759
Fish	460	40	500	8.000000
Reptile	292	20	312	6.410256
Nonvascular Plant	1312	20	1332	1.501502
Vascular Plant	16864	184	17048	1.079305

Table 7: Chi-squared test and Fisher's and Barnard's exact test results

In [87]: # iterate through "protected_data" and run "calculate_exact_tests()" for each 2x2 contingency table (not for AA; if there is AB, exclude BA)

```
# create empty lists to store categories under Null and Alternative hypotheses
  null_hypothesis_results = []
alternative_hypothesis_results = []
  iteration = 0
   # calculate the total number of iterations
  total_iterations = (len(protected_data) * (len(protected_data) - 1)) // 2
  # set the desired p-value threshold; 5% can be considered as the universal standard
# Divided by "total_iterations" to include the Bonferroni correction
# Expected output is 21 calculations; Checks out!
  p_value = 0.05 / total_iterations
   # loops that ensure the right amount of combinations for the contingency tables
  for i in range(len(protected_data)):
    not_protected_1 = protected_data.iloc[i]["not_protected"]
    protected_1 = protected_data.iloc[i]["protected"]
    category_1 = protected_data.index[i]
         for j in range(i+1, len(protected_data)):
    not_protected_2 = protected_data.iloc[j]["not_protected"]
    protected_2 = protected_data.iloc[j]["protected"]
                category_2 = protected_data.index[j]
                result = calculate_exact_tests([[not_protected_1, protected_1], [not_protected_2, protected_2]])
                \# compare the p-value to the given threshold and store everything \# force comparison with the most rigorous test (Barnards)
                if result["p-value"][2] < p_value:
    alternative_hypothesis_results.append({</pre>
                              "Category 1": category_1,
"Category 2": category_2,
                             "Category 2": category_2,
"Chi2 statistic": result["Statistic"][0],
"Chi2 p-value": result["p-value"][0],
"Fisher statistic": result["Statistic"][1],
"Fisher p-value": result["p-value"][1],
"Barnard statistic": result["Statistic"][2],
"Barnard p-value": result["p-value"][2],
"Hypothesis": "Alternative"
                      })
                else:
                       null_hypothesis_results.append({
                              "Category 1": category_1,
"Category 2": category_2,
                              "Chi2 statistic": result["Statistic"][0],
"Chi2 p-value": result["p-value"][0],
                             Fisher statistic": result["Statistic"][1],
"Fisher p-value": result["P-value"][1],
"Barnard statistic": result["Statistic"][2],
"Barnard p-value": result["p-value"][2],
"Hypothesis": "Null"
                      })
                print(f"Iteration {iteration}/{total_iterations} completed!")
   # create DataFrames to categorize the results
  null_hypothesis_df = pd.DataFrame(null_hypothesis_results)
  alternative_hypothesis_results = pd.DataFrame(alternative_hypothesis_results)
  # merge null and alternative hypothesis DataFrames
  exact_tests_results = pd.concat([null_hypothesis_df, alternative_hypothesis_results], ignore_index=True)
  print("Calculations successfully completed!")
Iteration 1/21 completed!
Iteration 2/21 completed!
Iteration 3/21 completed!
Iteration 4/21 completed!
Iteration 5/21 completed!
Iteration 6/21 completed!
Iteration 7/21 completed!
Iteration 8/21 completed!
Iteration 9/21 completed!
Iteration 10/21 completed!
Iteration 11/21 completed!
Iteration 12/21 completed!
Iteration 13/21 completed!
Iteration 14/21 completed!
Iteration 15/21 completed!
Iteration 16/21 completed!
Iteration 17/21 completed!
Iteration 18/21 completed!
Iteration 19/21 completed!
Iteration 20/21 completed!
Iteration 21/21 completed!
Calculations successfully completed!
```

Chi-squared test and Fisher's and Barnard's exact test results (Significance level: 0.002381) Category 1 Category 2 Chi2 statistic Chi2 p-value Fisher statistic Fisher p-value Barnard statistic Barnard p-value Hypothesis 0 Mammal Bird 0.970184 0.324634 0.883777 0.305849 -1 045225 0.298403 0.473148 1 Amphibian 11.127833 0.000850 0.000505 -3.431969 0.007760 Mammal 2 Bird Amphibian 8.793312 0.003023 0.535370 0.001834 -3.051553 0.002696 Null 3 Amphibian Fish 0.092023 0.761621 0.894410 0.697297 -0.433361 0.676771 Null 4 1.010914 0.314684 0.704501 0.293558 -1.155637 0.297246 Null Amphibian Reptile 5 Fish Reptile 0.496226 0.481163 0.787671 0.490707 -0.842331 0.409732 Null 6 Reptile Nonvascular Plant 23.632806 7 0.000000 Reptile Vascular Plant 70.459002 0.159298 0.000000 -8.659054 0.004388 Null Nonvascular Plant Vascular Plant 1.640202 0.200298 0.715750 0.172131 -1.416483 0.156892 8 Fish 9 19.981578 0.000008 0.423188 0.000004 -4.556220 Mammal 0.000146 Alternative 10 Mammal Reptile 19.696214 0.000009 0.333333 0.000003 -4.536699 0.000152 Alternative 0.074187 11 Mammal Nonvascular Plant 171.365605 0.000000 0.000000 -13.182738 0.000000 Alternative 12 Mammal Vascular Plant 1014.476132 0.000000 0.053099 0.000000 -31.999062 0.000000 Alternative 13 Fish 17.484891 0.000029 0.478841 0.000009 -4.254011 0.000929 0.377169 14 Bird Reptile 17.058216 0.000036 0.000007 -4.217667 0.000993 Alternative 15 Bird Nonvascular Plant 0.000000 0.083943 0.000000 -13.157489 171.546322 0.000000 16 Vascular Plant 1434.875551 0.000000 0.060082 0.000000 -37.955579 0.000000 Bird Alternative 17 Amphibian Nonvascular Plant 46.348633 0.000000 0.156794 0.000000 -6.994033 0.000248 Alternative 18 Vascular Plant 149.383640 0.000000 0.112226 0.000000 -12.480747 0.000041 Amphibian Alternative

19

20

Fish Nonvascular Plant

Vascular Plant

46.433369

179.167196

0.000000

0.000000

0.175305

0.125474

Figure 14: Testing for Significant Differences between Species in their Conservation Status using Barnard's exact test: A Distribution of p-values

-6.961541

-13.587424

0.000320 Alternative

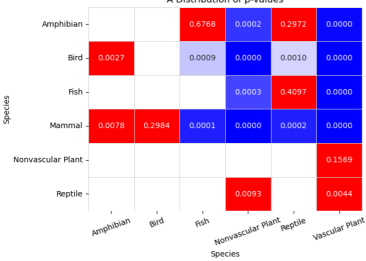
0.000000

0.000000

0.000000

```
In [89]: # select data for a heatmap
          heatmap_data = exact_tests_results[["Category 1", "Category 2", "Barnard p-value"]].copy()
          # pivot the data to create the heatmap data DataFrame
          heatmap_data = heatmap_data.pivot(index="Category 1", columns="Category 2", values="Barnard p-value")
         heatmap = sns.heatmap(heatmap_data, annot=True, fmt=".4f", cmap="bwr", cbar=False, vmin=0, vmax=p_value, linewidths=0.5, linecolor="lightgray")
          # add annotation below the araph for the hypotheses
         plt.annotate("Red corresponds to the Null Hypothesis\nBlue corresponds to the rejection of the Null Hypothesis",
                        xy=(0.8, -0.4),
xycoords="axes fraction",
                        ha="center
                        fontsize=12)
         # add annotation below the graph for significance level
plt.annotate("Significance level: 0.00238",
                        xycoords="axes fraction",
                        fontsize=12)
          # set the title, x-label, and v-label
         plt.title("Testing for Significant Differences between Species in their Conservation Status using Barnard's exact test:\nA Distribution of p-values")
          plt.xlabel("Species")
          plt.ylabel("Species"
          # rotate the x-axis tick labels
         heatmap.set_xticklabels(heatmap.get_xticklabels(), rotation=20)
         plt.savefig("Figure14.png", dpi=300, bbox_inches="tight")
          # display the figure
         plt.show()
```

Testing for Significant Differences between Species in their Conservation Status using Barnard's exact test: A Distribution of p-values



Red corr Significance level: 0.00238 Blue correspond

Red corresponds to the Null Hypothesis Blue corresponds to the rejection of the Null Hypothesis

Conclusions

The analysis of biodiversity has been conducted in four USA national parks:

- 1. Yellowstone National Park
- 2. Yosemite National Park
- 3. Great Smoky Mountains National Park
- 4. Bryce National Park

Although the surface areas of the areas in question vary significantly (Surface Areas Comparison), there is a noticeable pattern in the number and distribution of observed species, as depicted in Figure 1. These findings reinforce the suspicions that emerged during the data cleaning process, indicating that this dataset is fictional. Additional evidence can be found in Table 2. It is highly improbable for four national parks with distinct surface areas and geographical locations to exhibit such a perfect distribution of species, both collectively and individually, in terms of their conservation status. While such patterns may be observed in places like zoos, even there, they are unlikely to occur. Nonetheless, we can still conduct analysis and gain insights from this dataset, which comprises information on 5541 unique species across seven categories:

- 1. Mammal
- 2. Bird
- Reptile
 Amphibian
- 5. Fish
- 6. Vascular Plant
- 7. Nonvascular Plant

The most observed species in each park is a Vascular Plant called Sedge (https://en.wikipedia.org/wiki/Cyperaceae). Five most observed species for every national park can be seen in Figure 1.

Figure 2 displays the top five most observed species for each of the seven categories, with the count of observations displayed atop each bar. Observations serve as one of the primary analysis factors in this dataset. Vascular plants hold an overwhelming lead with 77% of the total observations, as depicted in Figure 3. This distribution aligns with expectations, as plants exhibit the widest range, remain stationary, are easily identifiable, and possess a high species abundance.

Even with the perfect distribution of species within parks, a touch of realism emerges when examining the total observations per park, as evidenced by the data provided (Figure 4). Yellowstone National Park stands out with 44% of all observations, coinciding with its status as the largest park in terms of surface area. However, there is a notable discrepancy between Bryce and Great Smoky Mountains, which cannot be accounted for due to the lack of information.

The count and relative proportions of observations for each category in each national park can be observed in Figure 5. The distribution of observations follows an established pattern, with Vascular Plants and Yellowstone National Park having the highest number of observations. Due to the significant difference in total observations per category, a logarithmic scale is applied to enhance the readability of proportions between categories. To gain insights into the distribution of species within each category, one can refer to Figure 6, and consult Table 1 for a more comprehensive analysis.

The dataset also comprises five distinct conservation status groups for each species, which serve to indicate the degree to which a species is threatened or protected:

- 1. Species of Concern
- 2. Endangered
- 3. Threatened
- 4. In Recovery
- 5. Least Concern

Figure 7 provides an overview of the conservation status distribution, revealing that the "Least Concern" category encompasses more than 96% of the total species within the dataset. For a more visual type understanding of conservation status distributions, Figure 8 presents a detailed breakdown for each category, employing a logarithmic scale yet again to ensure clarity and ease of interpretation. For a comprehensive numerical analysis, refer to Table 3, which contains the same information regarding conservation status distributions. Additionally, Figure 13 provides **a heatmap visualization corresponding to the data presented in Table 3**. Table 4 presents tangible results in the form of percentages, offering another clear representation of the data. Species of concern, endangered species, and species that are threatened are then analyzed in more detail.

Figures 9, 10 and 11 present a summary of the conservation statuses for endangered, species of concern, and threatened species, respectively. Among the realm of endangered species, mammals occupy a prominent position as one of the most vulnerable groups. A significant proportion of species of concern belongs to the avian class. However, it is the fish species that predominate as the majority of the threatened species. In addition to each figure, there is a table highlighting the top ten species observed for their respective conservation status. Figure 12 displays the total number of observations for each conservation status, and it aligns with the previous conclusions. As most species fall under the category of "Least Concerned", they are also the most observed ones. This pattern continues in a predictable way and in conjunction with previous data.

A **diversity index** is a quantitative measure that reflects how many different types (such as species) there are in a dataset (a community), and that can simultaneously take into account the phylogenetic relations among the individuals distributed among those types, such as **richness**, **divergence** or **evenness**. **These indices are statistical representations of biodiversity in different aspects**.

Table 5 presents the results for the Shannon-Weaver and Simpson's indices calculated for each national park. The findings indicate that all of the parks demonstrate substantial species richness and evenness in terms of the observed species. Additionally, there is a relatively low dominance of any particular species, indicating a balanced distribution of individuals among different species within the parks. These results collectively signify a high level of diversity among the species present in all four parks. Thus, it can be inferred that these parks exhibit notable biodiversity, characterized by a wide variety of species and a relatively equitable distribution of individuals among them.

There are several ways to determine associations between two categorical variables. In order to assess the relationship between protected and unproteced species categories, Chi-squared test, along with Fisher's, Barnard's, and Boschlo's exact tests, have been considered. The rationales for considering each test are discussed in section 5.4.2.

To ensure accurate assessment of the results, the significance level, also known as alpha (α), needs to be corrected. Therefore, the p-value is adjusted using the Bonferroni correction (5.4.2). Subsequently, Python and its modules are utilized to perform the aforementioned tests on several 2x2 contingency tables derived from Table 6. Boschloo's exact test is not considered in this analysis due to its use for ordinal categories.

Table 7 contains the results for each category pair, including the corresponding statistic and p-value for each test. Additionally, there is a "Hypothesis" column indicating whether the Null Hypothesis is accepted or rejected. In case of rejection, it states "Alternative". The significance level was compared to the most rigorous test, which is the Barnard's exact test. To provide a more visual representation of these numbers, Figure 14 illustrates a heatmap showcasing all sensible combinations of protected and unproteced species categories.

What can be gathered from these results is that mammals and birds are the most endangered categories because they account for the highest percentage of all protected species, followed by fish and amphibians. Some protected and unprotected categories show statistically significant differences, while others do not.

In order to gain a more comprehensive understanding of the results obtained from the tests conducted on significant differences between species in their conservation status using Barnard's exact test, it is essential to obtain additional data and perform more thorough analysis. The distribution of p-values obtained thus far indicates the presence of some potential distinctions, but to draw reliable conclusions and make informed decisions, further data collection and refined statistical examination are imperative. By incorporating a broader range of data and implementing robust analytical techniques, we can obtain a deeper comprehension of the variations in conservation status among species.

In order to safeguard biodiversity, it is imperative to allocate future resources towards the conservation and protection of endangered species. Among these, special emphasis should be placed on the Gray Wolf species, which faces significant threats to its survival.

Furthermore, it is essential to investigate the factors contributing to the substantial number of bird species falling under the "Species of Concern" conservation status, as well as the fish species classified as "Threatened". Several **potential risk factors** may be influencing these conservation statuses, including:

- Inadequate waste management
- Detrimental impact of wildfires
- Unsustainable hunting practices
- · Rapid pace of modernization

To address these issues effectively, a comprehensive approach is required. Adequate funding should be allocated towards habitat preservation, restoration, and management, with a focus on protecting critical ecosystems for these species. Collaborative efforts involving governmental organizations, conservation agencies, and local communities must be encouraged to develop and implement conservation strategies.

Moreover, **public awareness campaigns should be initiated to educate communities** about the importance of conserving these species and the ecosystems they inhabit. Promoting sustainable practices, such as responsible waste management and wildlife-friendly land use, can help mitigate the threats faced by these species.

Additionally, **robust legislation and enforcement mechanisms** should be put in place to **deter illegal activities**, including poaching and the trade of endangered species. Strengthening existing laws and regulations will ensure that the protection of these species remains a top priority.

In conclusion, directing future resources towards the preservation of endangered species, particularly the Gray Wolf, is of paramount importance. Simultaneously, a comprehensive assessment of the factors contributing to the increased number of bird species under the "Species of Concern" category and fish species in the "Threatened" conservation status is crucial. By addressing potential risk factors, implementing conservation measures, and raising public awareness, we can work towards safeguarding these species and their habitats for the benefit of present and future generations.