

# 5.2 Classification & Cladistics

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# 5.2.1 Classification System

# **Binomial System**

- The diversity of life on Earth is vast, and is known as global biodiversity
- The extent of global biodiversity is such that scientists can only **estimate** the total number of species present on Earth, and it is likely that there are **many species yet to be discovered**
- For biologists to make sense of the huge array of species, **organising them into logical groups** is essential
  - This process of putting organisms into groups is known as **classification**
  - The science of classification is known as **taxonomy**, and scientists working in the field of taxonomy are **taxonomists**
- Classifying an organism involves deciding which biological group, or taxon (plural taxa), it fits into best, and then naming it according to its taxon
- Historically an organism's biological group was determined on the basis of its observable characteristics, and today this information is combined with DNA sequence data for more accurate classification

### A Universal Naming System

- The biological system of naming used to name species according to their taxa is known as the binomial system
- This system is universal, ensuring that scientists around the world all use the same method of naming species
  - In the past, individual scientists decided on species names; names could be very **long**, and often one species could have **different names** in different parts of the world
- To ensure that all biologists know, and agree on, the criteria for naming species, regular meetings called **congresses** are held to discuss naming conventions
  - The first International Zoological Congress was held in 1889, during which taxonomists agreed on the rules that should be used for classifying and naming species
  - Congresses have since been held at regular intervals, with separate meetings for scientists who study different groups of organisms e.g. animals, plants, and fungi

### The Binomial System

- The **binomial naming system**, or system of nomenclature, was introduced by the Swedish taxonomist **Carl Linnaeus** in his 1758 book, Systema Naturae
- The system involves giving a species a **two-part name**, hence binomial
- Both parts of the name are in Latin, or a latinised version of a non-Latin word
  - e.g. Eriovixia gryffindori is a species of orb spider named after a famous school house
- The first part of the name is an organism's **genus**, and the second is its **species** name
  - E.g. the binomial name of a wolf is Canis lupus; wolves belong to the genus Canis, and the species lupus

# Your notes

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- There are several **conventions**, or rules, that should be used when writing binomial names
  - The genus should begin with a capital letter, and the species with a lower-case letter, e.g. the honey bee is Apis mellifera
  - When typed, binomial names should appear in italics, and when written by hand, they should be underlined e.g. a limpet is *Patella vulgata* when typed, or <u>Patella vulgata</u> by hand
  - The first time a binomial name is used in a text it should appear **in full**, e.g. wheat is *Triticum* aestivum, but the **genus name can from then on be abbreviated** to *T. aestivum*

# NOS: Cooperation and collaboration between groups of scientists; scientists use the binomial system to identify a species rather than the many different local names

- The work of scientists does not take place within the **confines of a laboratory**, or even a **country**, but can have implications for the work of other scientists around the world
- Because of the international nature of scientific research, it is essential that scientists are able to communicate with each other clearly about their work
- For scientific communication to be **effective** and **allow collaboration**, it is essential that scientific language enables scientists to be sure that they are **talking about the same thing**
- The **binomial naming system** is a good example of a system that enables scientists to communicate clearly with each other about living organisms
- The binomial system is essential because it ensures that scientists are all **talking about the same species** 
  - Different countries may have different common names for the same species e.g. In English, Felis domesticus is known as a cat, but around the world it is also known as kitte, maow, chat, kissa, bili, and gato
  - Different local or cultural names may arise even within a country e.g. in the UK alone the woodlouse, Onescus asellus, has more than 50 different local names that include woodpigs, cheesy bobs, woodywigs, chuggy pigs, and crunchy bats, and another country may have an equally long list of local, or cultural names for the same species
  - There are also examples of species where **one name might be used for different species** in different parts of the world, e.g. the robin in the US is a completely different species of bird to the robin in Europe
- There is nothing wrong with maintaining the use of different common names around the world, but when scientists discuss their work, the binomial system ensures that they are **communicating effectively** with each other

## Taxonomy

- Biological classification involves putting organisms into groups, or taxa (singular taxon)
- The taxa form a **hierarchy** 
  - A hierarchical system is one in which larger groups contain smaller groups with no overlap between groups
- The smallest taxon in the taxonomic hierarchy is **species**
- The species taxa are grouped within the next biggest taxon in the hierarchy, **genus** (plural **genera**)
- The genera are grouped within the next taxon, **family**, and so on until the biggest taxon, **domain**

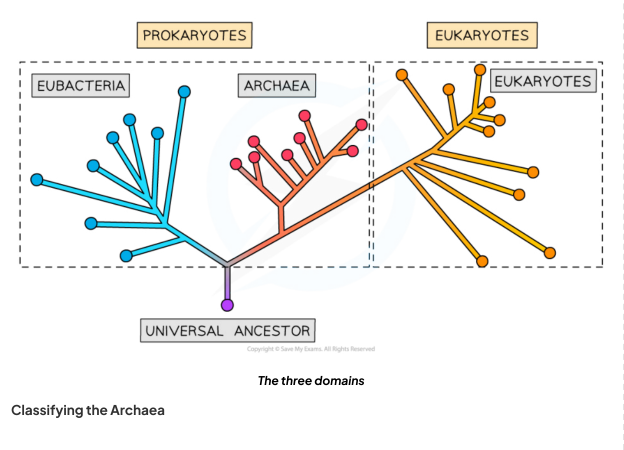
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# Domains

- Taxonomy is the practice of biological classification
  - Organisms are **grouped into taxa**, with the smallest taxon being species
- The science of taxonomy has frequently changed to match the latest discoveries about the features of organisms
  - Historically, the largest taxonomic groups were the **plant** and **animal kingdoms**, then **fungi** were discovered and added (incorrectly) to the plant kingdom
  - Microscopes led to the discovery of prokaryotes and eukaryotes, and the taxa were later divided into five kingdoms; plants, animals, fungi, protoctists, and prokaryotes
    - The protoctists are eukaryotic, primarily single-celled, organisms
  - RNA analysis has recently shown that there are two distinct groups of prokaryotes, leading to a shift in taxonomic thinking and the beginnings of the three domain system
- The largest taxonomic group is now known as a domain
- There are three domains, which are:
  - Archaea (prokaryotes)
  - Eubacteria (prokaryotes)
  - Eukaryotes (eukaryotes)





- The archaea were originally classified with the rest of the bacteria in one taxon due to **sharing several features** with them
  - Prokaryotic cell structure
  - Circular chromosome
  - Presence of a cell wall
  - 70S ribosomes
- Closer analysis of the archaea however showed that some of their features were distinct from the rest of the prokaryotes
  - Their cell walls are made of a different material from the cell walls of the rest of the bacteria
  - Their cell membranes are made of a distinct type of lipid
  - The **small subunit** of their **ribosomes** is **more similar to eukaryotic ribosomes** than to the ribosomes of the rest of the prokaryotes
- These discoveries led to a change in the classification of the archaea, which became their own domain

The Features of the Three Domains Comparison Table	
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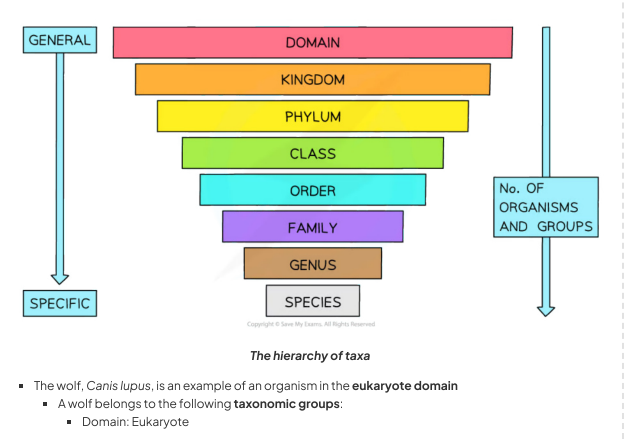
Feature	Archaea	Eubacteria	Eukaryotes
Cell type	Prokaryotic	Prokaryotic	Eukaryotic
Chromosome	Circular	Circular	Linear chromosomes + circular mtDNA and cpDNA
Cell membrane lipids	Glycerol-ether lipids	Glycerol-ester lipids	Glycerol-ester lipids
Ribosomes	<b>tibosomes</b> subunit is more similar to eukaryotic ribosomes		Larger 80S ribosomes in cytosol and 70S ribosomes in mitochondria and chloroplasts
Cell walls Always present (without peptidoglycan)		Always present (with peptidoglycan)	Sometimes present (without peptidoglycan)
Histones	Yes	No	Yes
Introns	Sometimes	Rarely	Yes

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# Your notes

# Hierarchy of Taxa for Eukaryotes

- Eukaryote is the domain of **all eukaryotes**, distinguishable from the Bacteria and the Archaea which are **both prokaryotic** domains
- Just like the other domains, the Eukaryote domain contains a hierarchy of taxa
- The taxonomic hierarchy contains the following taxonomic groups in descending order of size:
  - Domain
  - Kingdom
  - Phylum
  - Class
  - Order
  - Family
  - Genus
  - Species
- It can be good to use a mnemonic, or memory aid, to help you remember the different ranks in the taxonomic classification system
  - There are lots out there, or you can make up your own, but here's an example that you might find helpful:
    - Do Keep Ponds Clean Or Fish Get Sick
    - Domain, Kingdom, Phylum, Class, Order, Family, Genus, Species



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- Kingdom: Animalia
- Phylum: Chordata
- Class: Mammalia
- Order: Carnivora
- Family: Canidae
- Genus: Canis
- Species: lupus

• The flowering plant *Hibiscus rosa-sinensis* is another example of an organism in the eukaryote domain

- It belongs to the following **taxonomic groups**:
  - Domain: Eukaryote
  - Kingdom: Plantae
  - Phylum: Angiospermae
  - Class: Dicotyledonae
  - Order: Malvales
  - Family: Malvaceae
  - Genus: Hibiscus
  - Species: rosa-sinensis

### The Classification of the Wolf and the Hibiscus Plant Table

Taxonomic Rank	Wolf	Hibiscus
Domain	Eukaryote	Eukaryote
Kingdom	Animalia	Plantae
Phylum	Chordata	Angiospermae
Class	Mammalia	Dicotyledonae
Order	Carnivora	Malvales
Family	Candidae	Malvaceae
Genus	Canis	Hibiscus
Species lupus rosa-sinensis		rosd-sinensis

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# 5.2.2 Classifying Organisms

# **Examples of Classification**

- Biological classification involves putting organisms into groups, or taxa (singular taxon)
- The taxa form a **hierarchy** 
  - A hierarchical system is one in which larger groups contain smaller groups with no overlap between groups
- The taxonomic hierarchy contains the following taxonomic groups in descending order of size:
  - Domain
  - Kingdom
  - Phylum
  - Class
  - Order
  - Family
  - Genus
  - Species

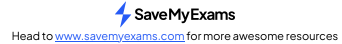
### **Examples of classification**

- The wolf, Canis lupus, is an example of an organism in the **animal kingdom** 
  - A wolf belongs to the following **taxa** 
    - Domain: Eukaryote
    - Kingdom: Animalia
    - Phylum: Chordata
    - Class: Mammalia
    - Order: Carnivora
    - Family: Canidae
    - Genus: Canis
    - Species: lupus
- The flowering plant *Hibiscus rosa-sinensis* is an example of an organism in the **plant kingdom** 
  - It belongs to the following taxa
    - Domain: Eukaryote
    - Kingdom: Plantae
    - Phylum: Angiospermae
    - Class: Dicotyledonae
    - Order: Malvales
    - Family: Malvaceae
    - Genus: Hibiscus
    - Species: rosa-sinensis

### The Classification of the Wolf and the Hibiscus Plant Table

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Taxonomic Rank	Wolf	Hibiscus
Domain	Eukaryote	Eukaryote
Kingdom	Animalia	Plantae
Phylum Chordata Angiosper		Angiospermae
Class	Mammalia	Dicotyledonae
Order Carnivora		Malvales
Family Canidae Malvaceae		Malvaceae
Genus Canis Hibiscus		Hibiscus
Species lupus rosa-sinensis		rosd-sinensis

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# Natural Classification

- The taxonomic classification system organises species into groups based on their evolutionary origins and relationships
- This means that all members of a taxonomic group **share a common ancestor** 
  - A common ancestor is one individual, or pair of individuals, from whom all individuals in a group have descended, e.g.
    - For a group of full siblings, the common ancestors will be their parents
    - For a group of cousins, the common ancestors will be their grandparents
- Grouping organisms into taxa that share a common ancestor is known as natural classification and a group of organisms classified in this way is known as a natural group
  - E.g. eukaryotic cells appear to have **only evolved once**, meaning that **one eukaryotic cell was the ancestor of all other eukaryotes**; the eukaryote domain is therefore a natural group
- It is possible to carry out artificial classification, grouping together organisms from different ancestors, e.g.
  - Dolphins and sharks could in theory be grouped together as they are both groups of aquatic animals that share a similar body shape, but they in fact **belong to different classes** 
    - Dolphins are **mammals** and sharks are **fish**
    - Their streamlined body shapes **evolved separately** rather than originating in one common ancestor
  - Cacti and euphorbia are two groups of desert plant recognisable by their spiny leaves and branching, succulent stems, and it would be reasonable to assume from their appearance that they are closely related to each other, but they **belong to different orders** of plant
    - Cacti are found in the deserts of the Americas, while euphorbia are found in Africa
    - They evolved separately, but **adapted to similar environments**, hence they have similar characteristics; this kind of evolution is known as **convergent evolution**
- It can be difficult to carry out accurate natural classification from observation alone, but the development of DNA sequencing technology means that natural classification on the basis of DNA sequence data has become possible

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SHARK
BUTTERFLY

Image: Constrained and constrained and points, and butterflies and bats, together on the second and constrained and points.

It would be possible to group both sharks and dolphins, and butterflies and bats, together on the basis of their shared characteristics, but this would not be a natural classification as they do not share recent common ancestors

### The advantages of natural classification

- Natural classification enables identification of unknown species
  - E.g. when a sample of insects is collected and the researcher doesn't recognise all of the species collected
  - Using the natural classification system an unknown species can be identified by logically working through the taxa
    - This means first identifying the domain, then the kingdom, then phylum etc., **narrowing down the options** until a species is identified
    - An identification method known as a **dichotomous key** works using this principle
- Natural classification enables predictions to be made about the characteristics of a species
  - If all of the members of a taxon descend from a **common ancestor** means that it is likely that the members will **share common characteristics**
  - This can be useful when e.g. searching for new drugs; if researchers know that one member of a plant genus produces a chemical that is useful in treating a disease, then it is likely that other members of that genus may produce the same chemical

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• This allows researchers looking for new drugs to focus their search rather than testing every single plant species



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# Reclassification

- There can be difficulties when trying to determine the ancestry of species based on observation of shared characteristics
  - This can lead to **artificial** rather than **natural classification** 
    - Convergent evolution can lead to groups of organisms **sharing similar characteristics** when they did not evolve from a shared common ancestor
- Advances in **DNA**, **RNA** and **protein sequencing** has allowed scientists to further investigate the relationships between species
  - This has revealed the true **ancestry** of taxa
- This sometimes leads to reclassification of taxa
  - Taxa may be **split** if taxonomists decide that they **do not descend from a common ancestor**
  - Taxa previously classified as separate may be grouped together if it is discovered that they do descend from a common ancestor
- An example of a taxon in which there has been a great deal of reclassification since the introduction of sequencing data is the Hominidea group to which humans belong
  - Humans were originally thought to have their own taxon separate from the rest of the great apes, but are now known to be part of the same family

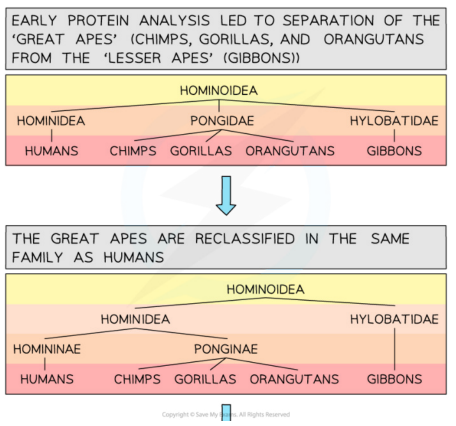
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	HOMINOI	DEA		
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HUMANS	CHIMPS	GORILLAS	ORANGUTANS	GIBBONS
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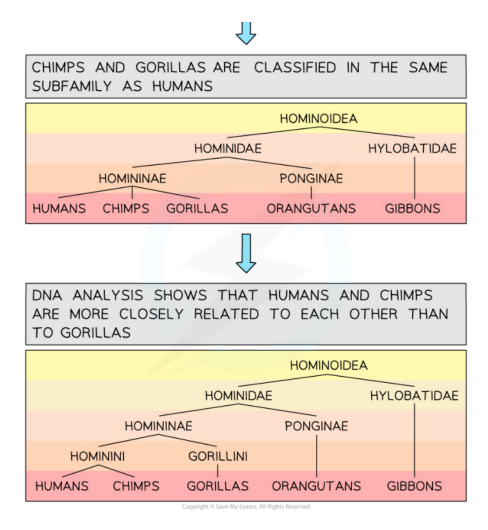
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The taxon containing humans has been frequently reclassified

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# 5.2.3 Plant Phyla Features

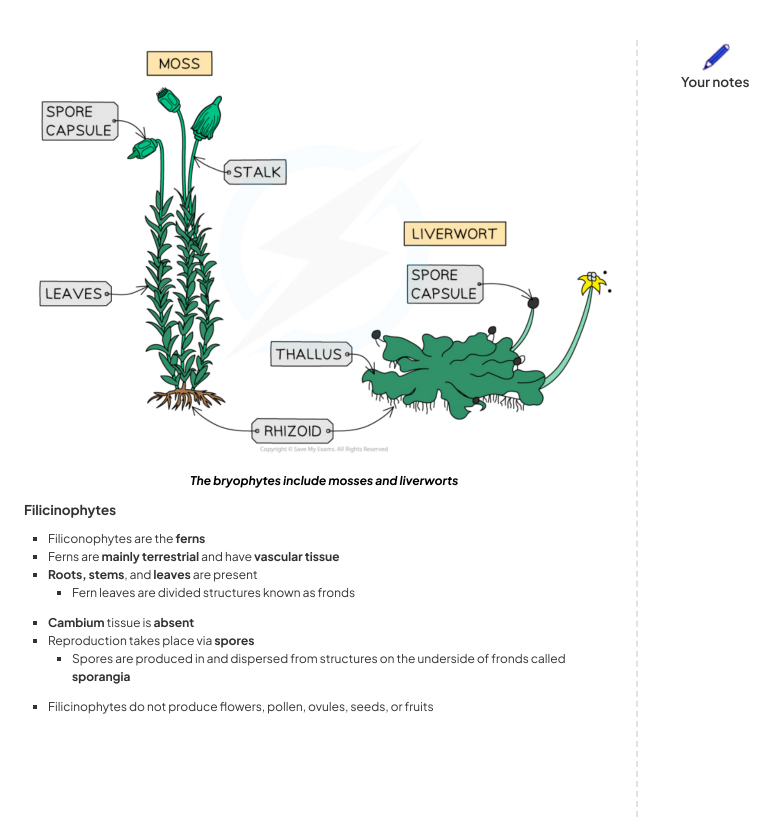
# **Plant Phyla Features**

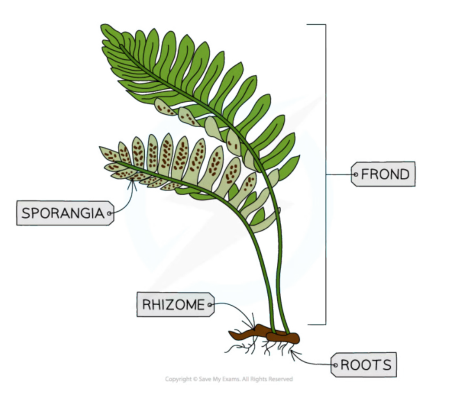
- All plants belong to the **plant kingdom**
- Within the plant kingdom are several **phyla** (singular **phylum**)
- Some plant phyla are small
  - E.g. the Glaucophytes contain only 70 species and the Gingkophytes contains only 1 living species
- There are four major plant phyla
  - Bryophytes (20 000 species)
  - Filicinophytes (10 000 species)
  - Coniferophytes (600 species)
  - Angiospermophytes (352 000 species)
- Each of the four main phyla can be identified by their characteristics

### **Bryophytes**

- The bryophytes include mosses, liverworts, and hornworts
- They are **small**, **terrestrial** plants
- They have no vascular tissue
- Cambium tissue is also absent
  - Cambium is a tissue containing stem cells that differentiate into xylem and phloem cells
- Reproduction takes place via **spores** 
  - Spores are produced in and dispersed from **spore capsules**
- Bryophytes do not produce flowers, pollen, ovules, seeds, or fruits
- No true roots are present, but hair-like structures called rhizoids anchor plants in the soil









#### The filicinophytes are the ferns

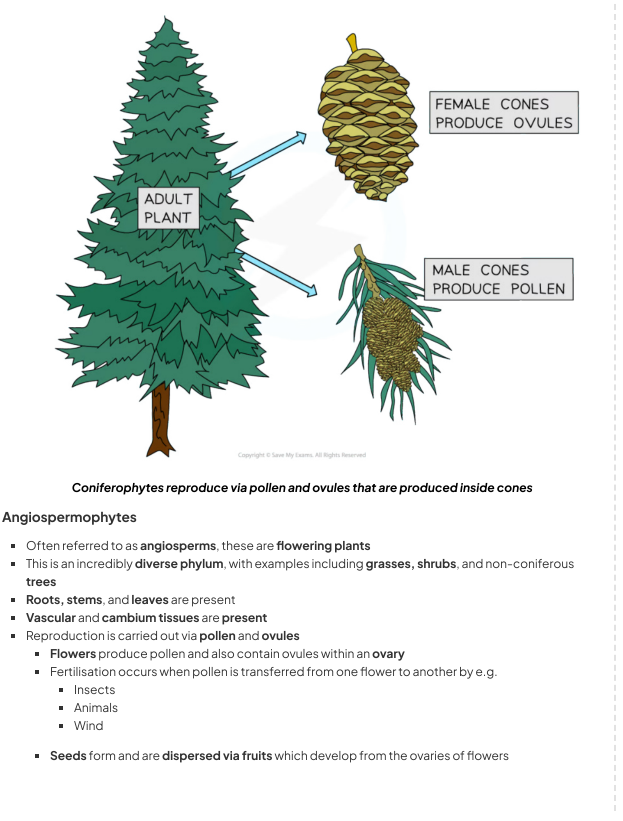
### Coniferophytes

- More commonly referred to as the **conifers**, the coniferophytes are usually **tall**, **straight**, **trees**
- Roots, stems, and leaves are present
- Vascular tissue is present, as well as cambium tissue
- **Reproduction** is carried out via pollen and **ovules** 
  - Pollen is produced in male cones and ovules are produced in female cones
     Pollen is the male gamete and ovules are the female gamete
  - **No flowers** are present
  - Both male and female cones are present on each tree
  - Pollen is carried from one cone to another by the wind in order for fertilisation to take place
- Seeds develop after fertilisation inside cones and are dispersed by falling to the ground or by animals
- Most coniferophytes are **evergreen** (i.e. they retain their leaves all year round)

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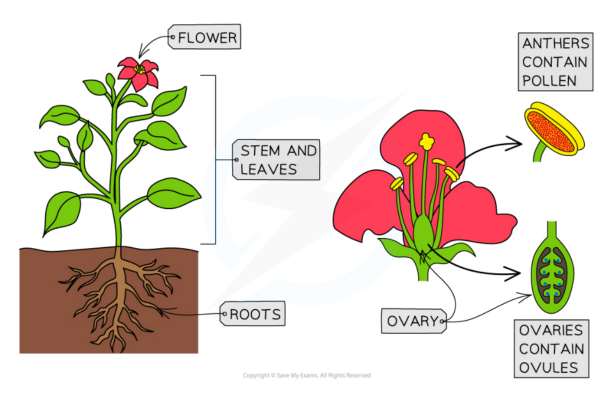
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#### Angiospermophytes reproduce via pollen and ovules produced in flowers

### Plant Phyla Features Table

Phylum	Plant structure under ground	Plant structure above ground	Vascular tissue	Mechanism of reproduction	Seeds	Fruits
Bryophyte	Rhizoid	Leaves or thallus	Absent	Spores in spore capsules	Absent	Absent
Filicinophyte	Roots	Leaves in the form of fronds	Present	Spores in spo- rangia	Absent	Absent
Coniferophyte	Roots	Stem and needle-like leaves	Present	Pollen and ovules in cones	Present in cones	Absent
Angiospermophyte	Roots	Stem and leaves	Present	Pollen and ovules in flow- ers	Present in ovdry	Present

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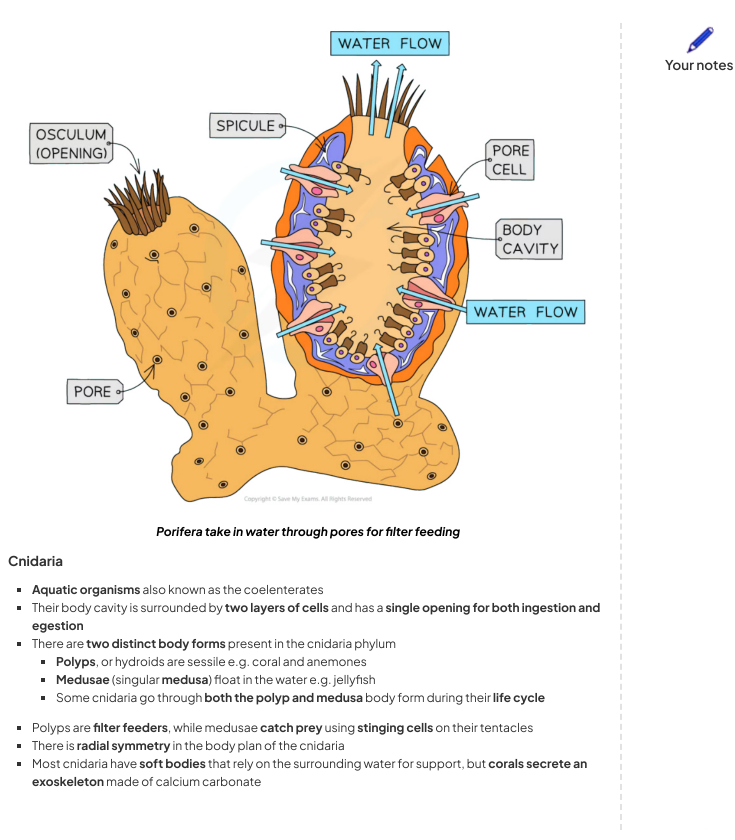
# 5.2.4 Animal Phyla Features

## **Animal Phyla Features**

- Animals are **multicellular**, **eukaryotic** organisms
- They are heterotrophs and most have a nervous system that enables responses to the surrounding environment
- There are over **30 animal phyla**, some of which are **vertebrates** and some of which are **invertebrates** 
  - Vertebrates have a backbone, or spinal column, while invertebrates do not
- Some of the major animal phyla include:
  - Porifera
  - Cnidaria
  - Platyhelminthes
  - Annelida
  - Mollusca
  - Arthropoda
  - Chordata

### Porifera

- Simple aquatic organisms consisting of a colony of cells
  - Commonly known as the **sponges**
  - Once thought to be plants
- Two layers of cells surround an internal cavity
  - They do not have a mouth for taking in food, or an anus for egestion
- Water is drawn in through pores in the outer layer of cells, before suspended particles from the water are filtered out and taken up by surrounding cells for food
  - Porifera are filter feeders
- There is **no symmetry** in the body plan of the porifera
- The structure of poriferans is supported by calcium or silica formations known as spicules

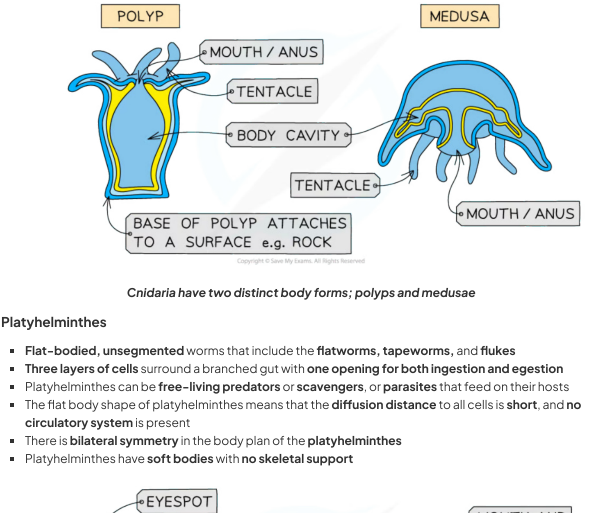


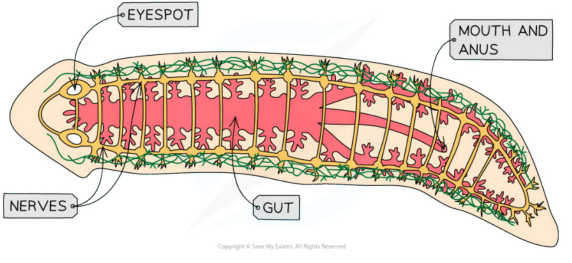
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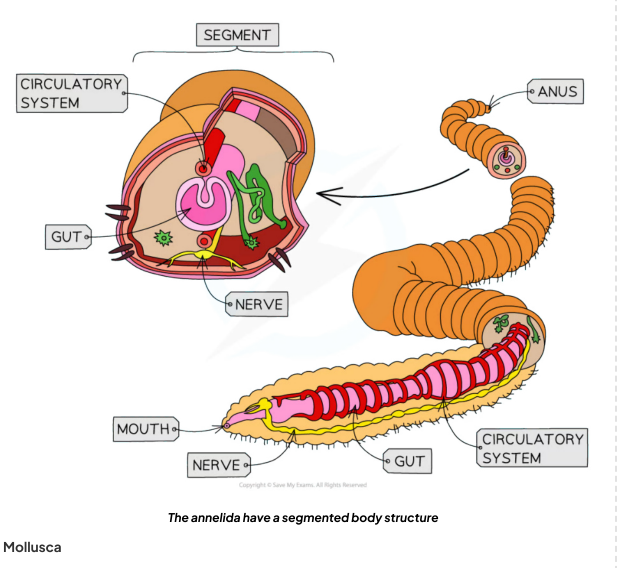
The body structure of a member of the phylum Platyhelminthes

Annelida

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- Round-bodied, segmented worms that include the earthworms and the leeches
- Each ring-shaped body segment contains a repetition of the structures of blood vessels and nerves
- The gut has separate openings for the mouth and anus
- Annelida can be free-living predators or scavengers, or parasites that feed on their hosts
- Annelida have a **closed circulatory system**
- There is **bilateral symmetry** in the body plan of the **annelida**
- Annelida have **soft bodies** which are supported by a **hydroskeleton** 
  - The hydroskeleton is an internal fluid-filled cavity called the coelom, against which the muscles of the body wall exert pressure



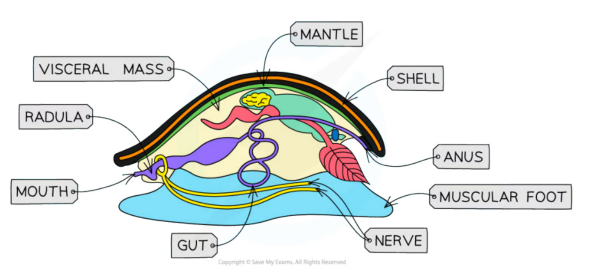
- A diverse group of animals that includes the **snails, slugs, mussels**, **limpets**, and **octopuses** 
  - It is the second-largest phylum (i.e. it contains a higher number of recorded species than any other phylum, except for Arthropoda)

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- Most molluscs are aquatic, with some terrestrial species
- The body structure contains a **muscular foot** and a **visceral mass** 
  - The foot is used for **motion** and/or **anchorage** e.g. as in limpets
  - The visceral mass is located above the foot and **contains the internal organs** (i.e. it is the soft, metabolic region that contains the digestive, excretory and reproductive organs)
- The **gut** has separate openings for the **mouth** and **anus**
- Mollusca have several feeding methods, including:
  - Filter feeding e.g. in bivalves such as mussels
  - Scraping at food sources such as leaves or algae on rocks using a rough, tongue-like organ called a radula e.g. snails and limpets
  - Active **predation** e.g. sea slugs
- Most mollusca have an **open circulatory system** in which the transport fluid bathes the internal organs
  - The cephalopods, which include squid and octopuses have a closed circulatory system
- There is **bilateral symmetry** in the body plan of the **mollusca**
- Mollusca have **soft bodies** that sometimes secrete a **calcium carbonate shell** from a mantle e.g. snails



The body structure of snails; members of phylum Mollusca

### Arthropoda

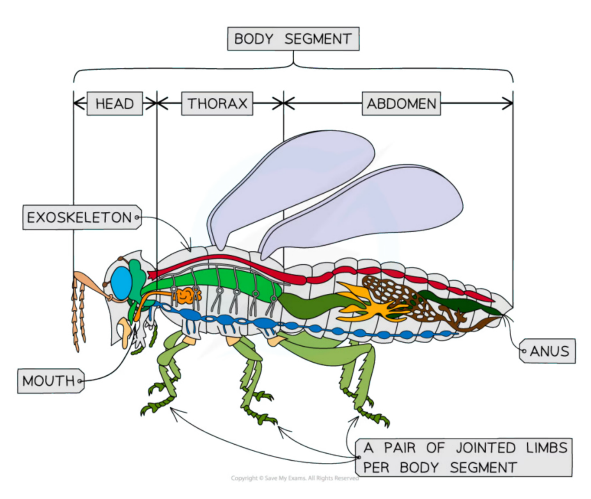
- Animals with segmented bodies, a hard exoskeleton, and jointed limbs
  - It is the **largest phylum** (i.e. it contains a higher number of recorded species than any other phylum)
  - It includes insects, arachnids, crustaceans, centipedes, and millipedes
- There is usually one pair of jointed limbs per body segment, though this is not always the case
- The **gut** has separate openings for the **mouth** and **anus**
- Arthropoda have an open circulatory system
- There is bilateral symmetry in the body plan of the arthropoda
- Arthropoda have a hard exoskeleton made of chitin

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• The exoskeleton is shed as the animal grows



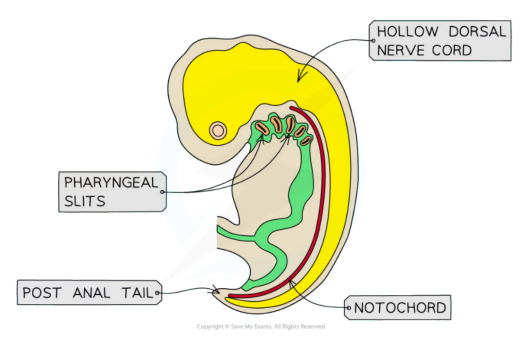
Insects are members of the phylum Arthropoda; they have three body segments, each with a pair of jointed limbs

### Chordata

- The phylum chordata includes all of the vertebrates, as well as some invertebrates such as the tunicates; commonly known as sea squirts
- To be classified as a chordate, an organism must have the following features at some stage during its development:
  - A rod-like structure along the **dorsal** length of the body known as a **notochord** 
    - Dorsal refers to the **back** of an organism
  - A hollow nerve cord located in the dorsal region
  - A series of slits behind the mouth known as **pharyngeal slits**
  - A tail that extends beyond the anus, known as a post-anal tail
- Some chordates only have these features during the **embryonic stages** of development, e.g.

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- In vertebrates, the notochord is replaced with the spinal column in fully developed individuals
- The dorsal hollow nerve cord develops into the brain and spinal cord in most species of chordate
- The pharyngeal slits are only retained into adulthood in some types of chordate e.g. in the gills of fish
- The post-anal tail has become a vestigial structure in the great apes
  - Vestigial structures are those that have lost their function, and are sometimes known as 'evolutionary leftovers'



### Human embryos have all of the features of phylum chordata

Phyla Features Table



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Phylum	Symmetry	Gut structure	Structural support	Other distinctive features
Porifera	None	None	Mineral structures called spicules	Filter water through pores on outer surface of body
Cnidaria	Radial	One opening for mouth and anus	Soft-bodied	Two distinct body forms; the polyp and the medusa
Platyhelminthes	Bilateral	One opening for mouth and anus	Soft-bodied	Unsegmented worms
Annelida	Bilateral	Separate openings for mouth and anus	Hydroskeleton	Segmented worms
Mollusca	Bilateral	Separate openings for mouth and anus	Soft bodied, but some also secrete a calcium carbonate shell	A muscular foot, and many have a radula
Arthropoda	Bilateral	Separate openings for mouth and anus	Hard exoskeleton	Segmented bodies and jointed legs
Chordata	Bilateral	Separate openings for mouth and anus Copyright © Save My Exams, All P	Internal skeleton	Must have a notochord, dorsal hollow nerve cord, pharyngeal slits, and a post—anal tail at some point during development

Your notes

# 오 Exam Tip

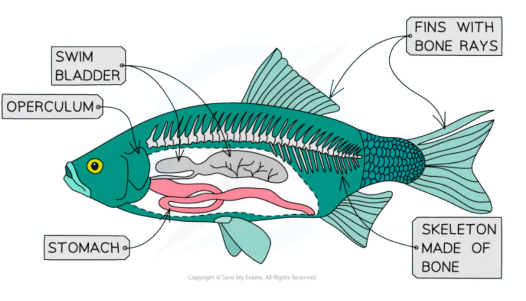
You do not need to be able to describe each of the above phyla in great detail, but be sure that you can state **how each phylum is distinct from the other phyla**, e.g. annelids are segmented worms while platyhelminthes are unsegmented worms, mollusca is the only phylum with a muscular foot, etc.

# **Vertebrate Features**

- The vertebrates are organisms that have a **spinal column** made up of **vertebrae**
- There are 5 major classes of vertebrate
  - Bony ray-finned fish (27 000 species)
    - There are other classes of fish, e.g. the bony lobe-finned fish (6 species), and the cartilaginous fish, whose skeletons are made of cartilage rather than bone (around 900 species)
  - Amphibians (6000 species)
  - Reptiles (9000 species)
  - Birds (10 000 species)
  - Mammals (5700 species)
- The 5 classes can be recognised by their features

### Bony ray-finned fish

- The ray-finned fish, also known as class Actinopterygii, have skeletons made of bone, and fins supported by rays, or spines, of bone
- The scales covering the bodies of the ray-finned fish vary in shape and composition
- Oxygen is gained via diffusion from the water into the gills, which are covered by a gill flap called the operculum
- During reproduction, eggs and sperm are released into the water and fertilisation takes place outside the body; this is external fertilisation
- Their entire life cycle is aquatic
- Body temperature is not regulated but changes with the surrounding environmental temperatures
- Buoyancy is regulated by inflating and deflating an internal air sac called a swim bladder



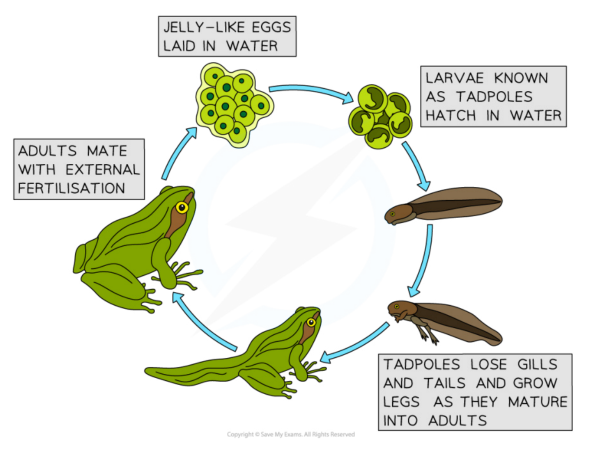
Ray-finned fish have a skeleton of bone, fins with bone rays, a swim bladder, and a gill operculum



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### Amphibians

- This class includes animals such as **frogs, toads**, and **salamanders**
- Amphibians have **soft**, **moist skin**
- Amphibian skin is permeable, enabling exchange of gases while under the water
  - Simple lungs are also present for gas exchange on land
- During reproduction, fertilisation is **external**
- Jelly-like eggs must be laid in water, and the larval stage of the life-cycle is aquatic e.g. frog tadpoles
  - Larvae such as tadpoles mature into an adult body form
  - Adult amphibians often leave the water and spend most of their time on land
  - There are a few exceptions to this; some frog and toad species can lay eggs on land which hatch into tiny adults rather than tadpoles
- Body temperature is **not regulated**



#### The amphibian life-cycle includes an aquatic larval stage, and a terrestrial adult stage

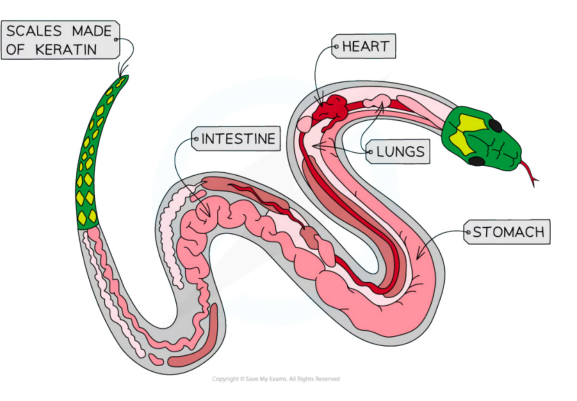
#### Reptiles

- This class includes animals such as **lizards, snakes**, and **turtles**
- Reptiles have impermeable, scaly skin made of keratin protein

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- Reptiles are mainly **terrestrial**, gaining their oxygen through highly folded surfaces in their **lungs**
- During reproduction, male sperm is deposited inside the female's body, so fertilisation is **internal**
- The eggs of reptiles are **soft-shelled**
- Reptile body temperature is **not regulated**



Reptiles have a body covering of scales, and breathe using highly folded lungs

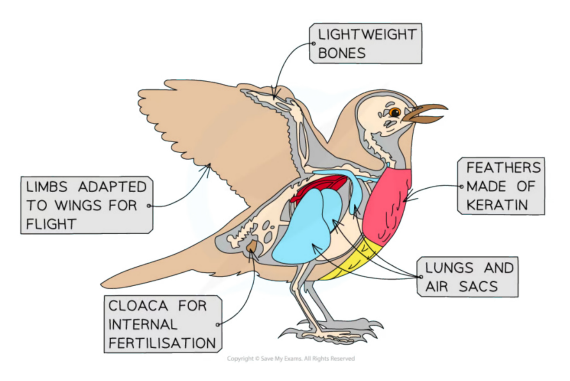
#### **Birds**

- This class includes animals such as chickens and ostriches
- Skin is covered with **feathers** made of keratin
- The front limbs of birds are modified into **wings**, and **skeletons are lightweight**, both adaptations for **flight**
- Birds gain oxygen via a pair of **lungs** and connected **air sacs**
- Fertilisation is internal
- The eggs of birds are **hard-shelled** 
  - Shells contain calcium carbonate
- Bird **body temperature is regulated**, so is often warmer than the surrounding environment

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#### Birds are usually highly adapted for flight and reproduce using internal fertilisation

#### Mammals

- This class includes animals such as elephants, humans, and whales
- Skin is covered with **hair** made of keratin
  - In some mammals body hair is lost in adulthood, e.g. in aquatic mammals
- Mammals gain oxygen via a pair of **lungs**
- Fertilisation is internal
- The young of mammals develops inside the body of the mother, and live young are born
  - There are some unusual exceptions to this e.g. the duck-billed platypus and the echidna
- Young mammals are fed on **milk** produced in **mammary glands**
- Mammal body temperature is regulated

Vertebrate Features Table

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Class	Skin covering	Gaining oxygen	Fertilisation	Reproduction	Body Temperature
Ray– finned fish	A variety of scale types	Gills	External	Eggs are laid and hatch in water. Hatchlings are fully developed	Unregulated
Amphibian	Soft, moist skin	Gills in larvae, then skin and lungs in adults	External	Jelly–like eggs are usually laid and hatch in water. Young are in the for of larvae which mature into adults	Unregulated
Reptile	Scaly skin of keratin	Lungs	Internal	Soft eggs are laid on land	Unregulated
Bird	Feathers of keratin	Lungs and air sacs	Internal	Hard eggs are laid on land	Regulated
Mammal	Hair of keratin	Lungs	Internal	Live young are born	Regulated

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# 5.2.5 Cladistics

### Clades

- The term **clade** can be defined as
  - A group of organisms that have all descended from a common ancestor
- Cladistics is the branch of science in which scientists put organisms into clades
  - It involves classification that is based on homologous characteristics rather than analogous characteristics
- Clades are formed on the basis of evolutionary relationships i.e. who is descended from which ancestor
- Note that while taxonomy is about classifying and then naming organisms, cladistics is about identifying evolutionary relationships between organisms
  - A **taxon** is a group of organisms that have been given a group name by taxonomists **on the basis on their shared features**
  - A clade is a group of organisms classified together on the basis of their shared descent from a common ancestor
- If taxonomy is carried out correctly then all of the members of a taxon should form a clade, but due to historical errors and the difficulties in distinguishing between true homologous characteristics and those that have come about by convergent evolution, this is not always the case
- Clades can include both living and extinct species
  - Some of the descendants of a common ancestor may have gone extinct
  - The common ancestor species itself may have gone extinct
- Clades can be large or small depending on the common ancestor being studied



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# Identifying Members of a Clade

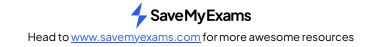
- In the past, scientists encountered many difficulties when trying to determine the evolutionary relationships between species
  - Using the physical features of species has **limitations** and can often lead to organisms being put into **groups that are not true clades**
  - This would mean that all of the organisms in a group are not descended from a common ancestor
     Some descendants might be missing
    - Some organisms might have been included that descend from a different ancestor
- Advances in sequencing technology have allowed scientists to further investigate the evolutionary relationships between species
- Sequence data that can be used to investigate evolutionary relationships can come from
  - DNA
  - mRNA
  - Amino acids in polypeptides
- Sequencing technology can determine the order of DNA bases, mRNA bases and amino acids
- For all types of sequence data, it can be said that the more similar the sequences, the

### more closely related the species are

- Two groups of organisms with very similar sequences have separated into separate species more recently than two groups with less similarity in their sequences
- Species that have been separated for longer have had a greater amount of time to accumulate mutations and changes to their DNA, mRNA and amino acid sequences
- Sequence analysis and comparison can be used to create family trees that show the evolutionary relationships between species

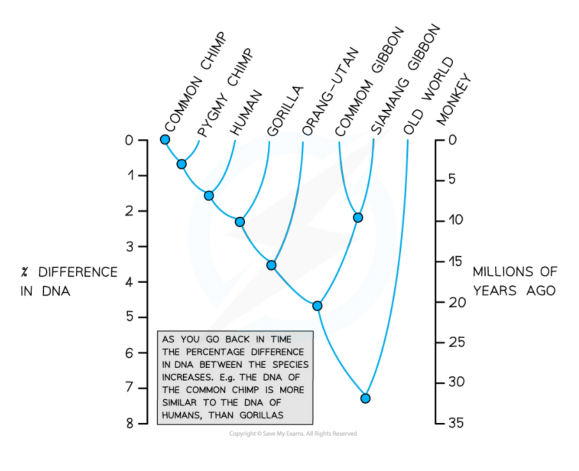
# 💽 Exam Tip

You may be wondering why you would use amino acids when you could look at DNA or mRNA; it is often easier to find and isolate proteins from cells than it is to isolate DNA or mRNA. However, DNA or mRNA analysis is often more powerful because genes for the same protein may have slightly different base sequences in different species.



# **Divergence from a Common Ancestor**

- The evolutionary relationships between species can be determined by analysing sequence data from e.g. DNA, mRNA, or amino acids in polypeptides
- The number of differences between sets of sequence data provides information on how closely related two species are
  - The more differences there are between the sequences, the longer ago the species diverged, and vice versa
- The differences between sequence data can also be used to produce a quantitative estimate for how long ago two species diverged from each other
  - Differences in sequence data come about due to mutations in the DNA
    - Evidence suggests that mutations occur at a constant rate
    - This means that the number of mutations that have occurred gives an indication of the amount of time that has passed since two species diverged
      - Scientists refer to the constant rate of mutation as the **molecular clock**
- Analysing the differences in sequence data allows evolutionary biologists to determine the order in which different species diverged from a common ancestor, and therefore how closely related species are



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Differences in DNA sequence data show how much time has passed since species diverged from each other, enabling the relationships between species to be established

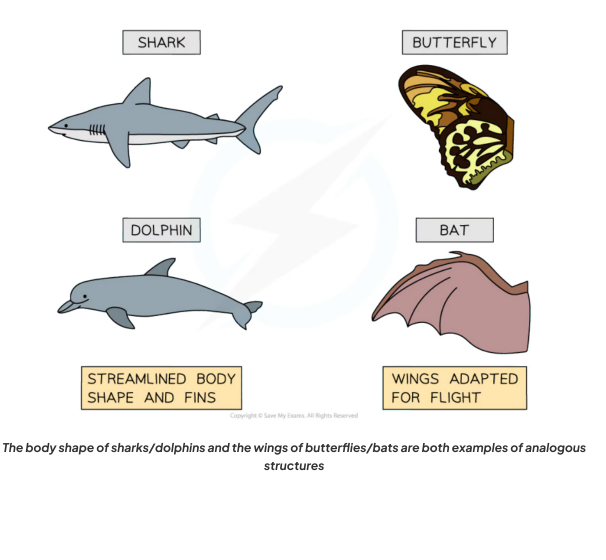


## Analogous & Homologous Traits

- Homologous traits can be defined as
  - Characteristics that may differ in form and function in different species but that have shared evolutionary origins
- Homologous traits, or characteristics, indicate common ancestry, and are useful for classifying organisms into true clades
  - An example of a homologous characteristic is the **pentadactyl limb**; limbs in different species of animal differ significantly in their shape and role, but similarities in overall structure indicate common ancestry
- The difficulty with using homologous traits in classification is that it is not always obvious whether characteristics are **homologous** or **analogous**
- Analogous traits can be defined as
  - Characteristics with the same function but which do not share an evolutionary origin
- Such characteristics have evolved independently of each other from different ancestors, enabling
  organisms to adapt to similar environments
  - This is known as **convergent evolution**
- Analogous characteristics **look similar**, hence the danger of confusing them with homologous characteristics
- Classifying organisms on the basis of analogous characteristics will **not produce an accurate clade** 
  - This has led to **errors** of classification in the past
  - For this reason, **sequencing data** is now used for classification instead of observable characteristics



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### Cladograms

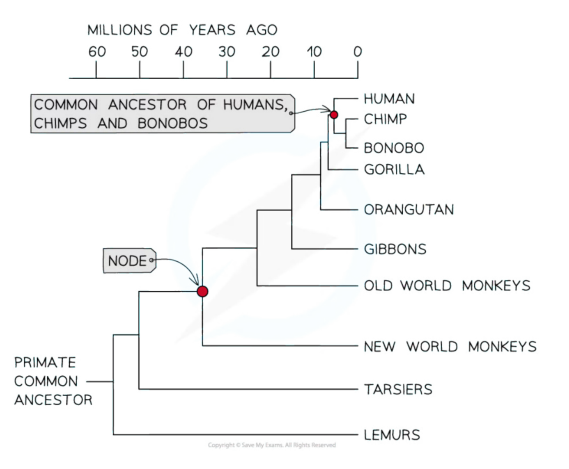
- Evolutionary relationships between species can be represented visually using a diagram called a cladogram
- Cladograms are evolutionary trees that show probable order of divergence from ancestral species and therefore probable relationships between species
  - The point at which two branches separate is known as a **node**
  - Nodes represent common ancestor species
- The information used to build cladograms most often comes from **sequence data** due to difficulties in the use of observable characteristics
  - It can be difficult to be sure whether observable traits are **homologous** or **analogous**
- Sequence data can provide information about how different species are from each other, as well as how much time has passed since divergence from a common ancestor took place
  - The constant rate at which mutations accumulate can be used as a **molecular clock**
- Computers use the information from sequence data to build the most likely cladogram
  - This is done using the principle of **parsimony**, which states that the simplest explanation is preferred
    - The computer builds the shortest possible cladogram with the smallest number of divergence events to fit the available data
  - We say that cladograms show the **most probable** divergence times and relationships rather than providing **definite** conclusions

#### Cladograms that include humans and other primates

- Analysis of sequence data for humans and other primate groups show that humans are most closely related to chimps and bonobos, and that the next closest relative is the gorilla
- Humans are thought to have diverged from chimps and bonobos between 5–7 million years ago

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A cladogram showing humans and other primates

## 5.2.6 Reclassification

### **Cladistics & Reclassification**

- The development of **DNA sequencing technology** means that classification can now be carried out on the basis of **evolutionary relationship** 
  - Organisms classified in this way are grouped into clades
- Historically, organisms would have been classified on the basis of observed traits, which often led to
  organisms being classified into groups that were not true clades
  - This resulted from analogous characteristics being incorrectly thought to be homologous
- The use of DNA sequencing to classify organisms has led to discoveries of classification errors and the need to **reclassify** organisms
  - Some species have been reclassified into **different groups** of organisms
  - Some groups of organisms have been **split**
  - Some groups have been merged
- Classifying organisms correctly according to their clade is important to ensure that groups of organisms are close evolutionary relatives
  - This means that the characteristics of one group member can be **predicted** on the basis of the characteristics of another

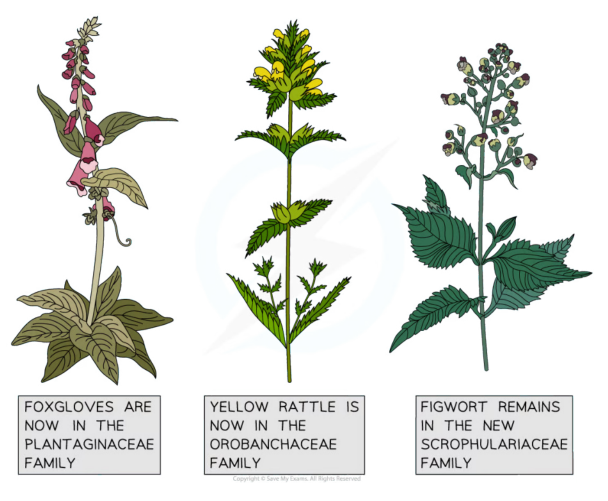


## **Example of Reclassification**

### Reclassification of the figwort family

- The figwort (pronounced 'figwurt') family, also known as the Scrophulariaceae, was at one point the 8th largest family in the angiosperm phylum
- When the group was originally classified in the late 1700s it contained 16 genera, which later expanded to 275 genera
  - The classification of the figworts was based on **observable traits** such as a tube-shaped flower structure
  - Examples of members of the original figwort family include foxgloves and yellow rattle
- When **DNA sequence analysis** began, plant scientists discovered that the shared features of the figwort family were in fact **analogous** and not evidence of **shared ancestry** 
  - Three chloroplast **genes** were analysed
  - It was discovered that the original figwort family was not a true clade
  - The figworts were found to contain **several separate plant families** 
    - New families were created
    - Several genera were moved into other existing families
    - The remaining genera were grouped together with two previously missed genera to form the **new figwort family**, still known as the Scrophulariaceae
- The new figwort family is **less than half** its original size and is only the 36th largest in the angiosperms, where it had previously been 8th

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#### DNA sequencing data led to the reclassification of many species within the original figwort family

NOS: Falsification of theories with one theory being superseded by another; plant families have been reclassified as a result of evidence from cladistics

- A theory is an explanation of observed phenomena that is supported by evidence
- This means that when **new evidence** is found that **no longer supports the theory**, it needs to be **changed** to take the new evidence into account
- Reclassification of organisms on the basis of DNA sequencing data is a good example of this
  - Scientists theorise that a group of plants should be classified together on the basis of existing evidence; historically this would have been the evidence of observable traits such as flower shape
  - New evidence from DNA sequence analysis shows that existing plant groups are not true clades, and so are not descendants of a common ancestor
  - The historical classification **theory has therefore been falsified** and needs to be changed to take the new evidence into account
  - New evidence from DNA and computer analysis is used to calculate the most likely plant evolutionary relationships, and this evidence is used to form a new theory regarding the

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classification of plants

• Analysis of new DNA data continues all the time, and if **evidence** is found that doesn't fit with current classification theories then more **falsification** and reclassification will take place



## 5.2.7 Skills: Classification & Cladistics

## **Dichotomous Keys**

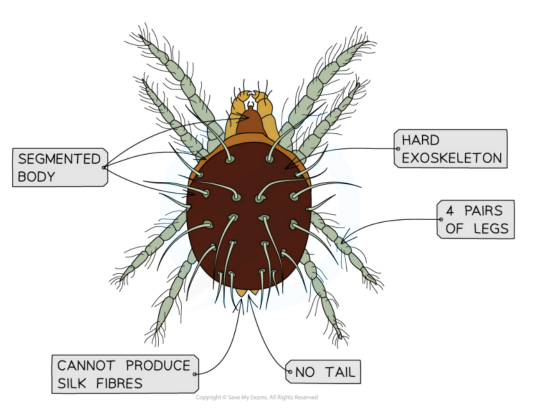
- For anyone who doesn't specialise in studying a particular group of organisms, it can be very **difficult** to identify, or ID, a species when working in the field
  - Correctly identifying species may be important for a researcher studying biodiversity or looking at the impacts of a changing environment on a community of organisms
- Someone seeking to identify species with which they are not already familiar may use a tool known as a **dichotomous key**
- A dichotomous key contains a series of **paired statements** 
  - The term 'dichotomous' refers to these pairs of statements
  - An example of such a pair of statements might read:
    - The organism shows radial symmetry
    - The organism shows bilateral symmetry
  - Or:
    - The organism has one pair of wings
    - The organism has two pairs of wings
- To work through a dichotomous key, you start with the first pair of statements and apply them to the unknown species; one statement will be clearly false, while the other will be a correct description of the species
- The correct statement leads to **another pair of statements**, and so on until the final correct statement leads to the name of the species



### Worked example

Use the dichotomous key provided to identify the type of organism below

The organism has an exoskeleton, a segmented body, and 4 pairs of legs. It has no tail and cannot produce silk





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1	Hard exoskeleton absent Hard exoskeleton present	
2	Hard shell absent Hard shell present	
3	3 pairs of legs 4 pairs of legs	
4	Tail absent Tail present	6 Scorpion
5	Body colour uniform Body striped black and yellow	
6	Silk producing organ present Silk producing organ absent	

#### Step 1: Decide which of the first pair of statements applies

The organism has an exoskeleton, so we can ignore the first statement in pair 1 and move onto the third pair of statements as instructed in the key

#### Step 2: Decide which of the third pair of statements applies

The organism has 4 pairs of legs, so we move on to the fourth pair of statements as instructed

#### Step 3: Decide which of the fourth pair of statements applies

The organism has no tail, so we know that it is not a scorpion, and we move onto the sixth pair of statements as instructed

#### Step 4: Decide which of the sixth pair of statements applies

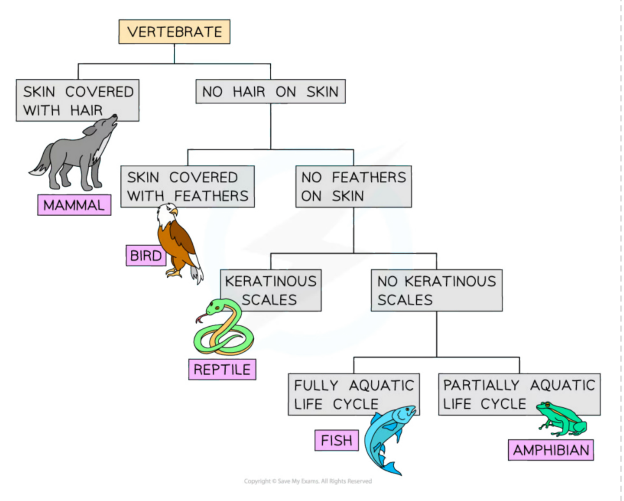
The organism cannot produce silk, so it must be a species of mite

#### Constructing a dichotomous key

• When constructing a key, the following should be considered

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- Each pair of statements should contain features that are **clearly identifiable** and **not subject to opinion** e.g. the organism has 3 pairs of legs, rather than e.g. the organism is small
  - A leg count gives an objective number, but size is relative and depends on what the species is being compared to
- Statements must be 'yes' or 'no' in style
- Each pair of statements should **divide the organisms being identified into two distinct groups**
- Each subsequent pair of statements should divide the organisms into **smaller and smaller groups**
- Each statement should be followed by either a **number** to continue the process of narrowing down the options, or should **name the organism** to which it applies



Dichotomous keys can also be represented using a branched diagram as shown here. This visual representation can be easier to understand, but it is limited in the number of organisms it can include

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### 😧 Exam Tip

Get some practice at using a dichotomous key by identifying the remaining organisms featured in the key, all of which are described below. Note: they are all quite easily recognisable animals so you should be able to tell whether you have used the key correctly or not.

- 1. The organism has an exoskeleton, a segmented body, and 4 pairs of legs. It has a tail with a sting
- 2. The organism has an exoskeleton, a segmented body, and 3 pairs of legs. Its body is uniform in colour
- 3. The organism has an exoskeleton, a segmented body, and 3 pairs of legs. It's body is striped black and yellow
- 4. The organism is soft bodied with a muscular foot and a hard shell
- 5. The organism has an exoskeleton, a segmented body, and 4 pairs of legs. It has no tail but can produce silk fibres from a specialised organ on its abdomen
- 6. The organism is soft bodied with a muscular foot and no hard shell

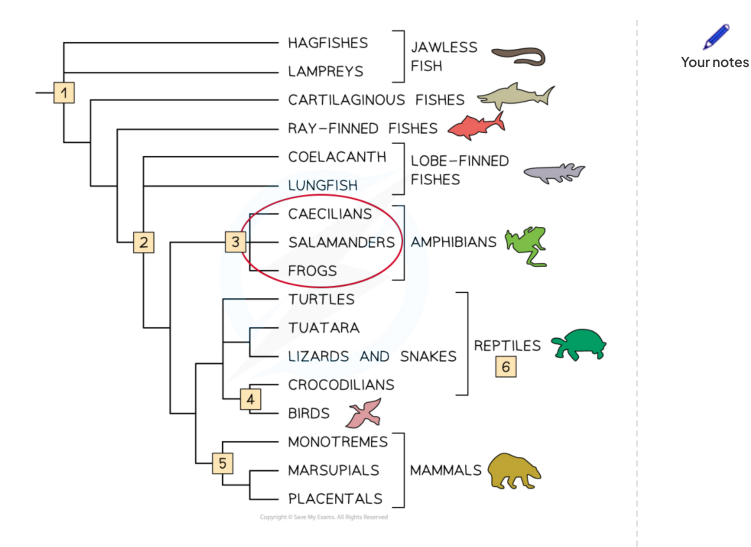
## Analysis of Cladograms

- Evolutionary relationships between species can be represented visually using a diagram called a cladogram
- Cladograms are evolutionary trees that show probable order of divergence from ancestral species and therefore probable relationships between species
- Analysis of a cladogram can provide several pieces of information
  - The point at which two branches separate is known as a node, and represents common ancestor species
    - A node **immediately adjacent** to a pair of clades indicates that these two clades share a **recent common ancestor**
    - This shows that the two clades are more closely related to each other than they are to any other clade in the cladogram
    - If several nodes need to be traced back before two clades can be joined, this indicates a more distant relationship between two clades
  - Cladograms sometimes show numbers along the branches; these indicate the number of base or amino acid changes that have occurred between one node and the next or between a node and an emerging clade or species
    - The constant rate at which mutations accumulate means that these numbers can be used as a **molecular clock** to calculate how much time has passed
  - Some cladograms have a time scale to show how many millions have years have passed
- Computers use the information from sequence data to build the most likely cladogram
  - This is done using the principle of parsimony which states that the simplest explanation is preferred
    - The computer builds the shortest possible cladogram with the smallest number of divergence events to fit the available data
- Cladograms provide the **most likely estimate** of the evolutionary progress of organisms
  - The reliability of a cladogram may vary depending on the amount of sequence data used to construct it
    - A cladogram based on the sequencing of one gene will be less reliable than a cladogram based on the sequencing of several genes
  - Cladograms are **subject to change** when new sequence data becomes available

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## 

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1	NODE REPRESENTING COMMON ANCESTOR OF ALL VERTEBRATES
2	FIVE NODES NEED TO BE TRACED BACK TO LINK PLACENTAL MAMMALS WITH COELACANTHS, INDICATING A MORE DISTANT RELATIONSHIP
3	THREE CLADES DIRECTLY LINKED TO ONE COMMON ANCESTOR INDICATES THAT THIS CLADOGRAM WAS BUILT WITHOUT ENOUGH DATA TO DETERMINE WHICH OF THESE THREE CLADES ARE MORE CLOSELY RELATED
4	ONLY ONE NODE NEEDS TO BE TRACED BACK TO LINK CROCODILIANS AND BIRDS, SO THEY ARE MORE CLOSELY RELATED TO EACH OTHER THAN TO ANY OTHER CLADE
5	NODE REPRESENTING COMMON ANCESTOR OF ALL MAMMALS
6	NOTE THAT THE REPTILES ARE NOT A TRUE CLADE, AS THEY DO NOT INCLUDE THE BIRDS, WHICH SHARE A COMMON ANCESTOR WITH THEM
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A cladogram with notes to indicate some of the conclusions that can be drawn from it. Note that this cladogram contains no numbers or time scale, so it does not show the number of base or amino acid changes that have occurred between one node and the next, or how much time has passed between nodes.

