

FACULTAD DE INGENIERÍA Y CIENCIAS EXACTAS DEPARTAMENTO DE BIOTECNOLOGÍA Y TECNOLOGÍA ALIMENTARIA UNIVERSIDAD ARGENTINA DE LA EMPRESA

Bioinformática

ANÁLISIS COMPUTACIONAL DE SECUENCIAS

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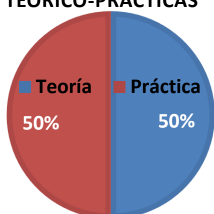
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Organización de la materia



CONTENIDO CLASES TEORICO-PRÁCTICAS





Parte I

- •Informe TP final I: Entrega por grupo, hasta 2 alumnos.
- Parcial Teórico-práctico
- Parte teórica (Escrito)
- •parte práctica (Ejercicios en computadora)

Parte II

- •Informe TP final II: Entrega por grupo, hasta 2 alumnos.
- Parcial Teórico-práctico
- Parte teórica (Escrito)
- parte práctica (Ejercicios en computadora)



Para aprobar el Final se debe aprobar la parte teórica y la parte práctica (>=4)

Recuperatorio de parcial I o II: Se recupera uno solo de los parciales

Misma modalidad que los parciales

Final anticipado Teórico y práctico

- Parte teórica (Escrito)
- Parte práctica (Escrito Interpretación de resultados)
- Para acceder a los finales deben estar aprobados los TPs y los parciales I y II

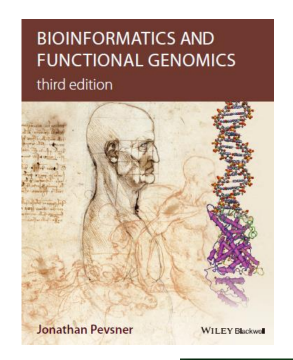


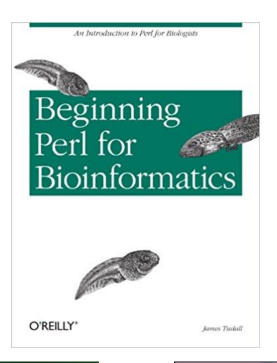
Para aprobar el Final se debe aprobar la parte teórica y la parte práctica (>=4)

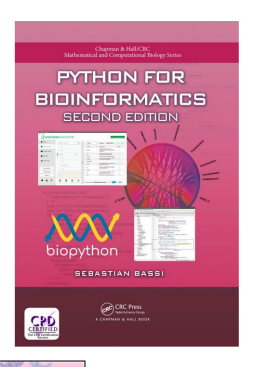


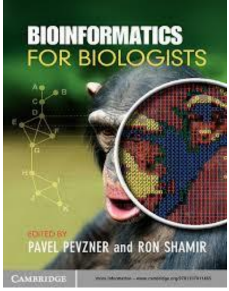
Para aprobar la materia de deben tener aprobados los 2 parciales, los 2 informes TP y el Examen Final

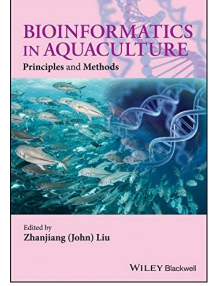
Lecturas recomendadas











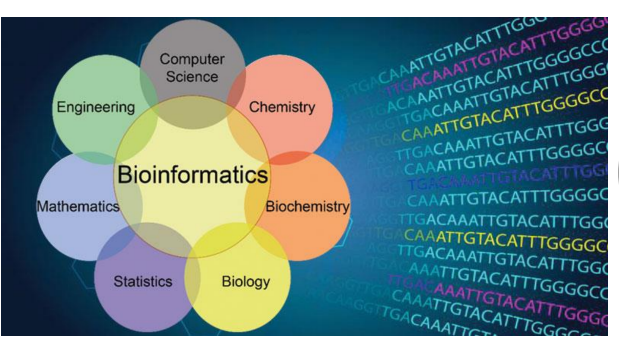
Objetivo de la materia

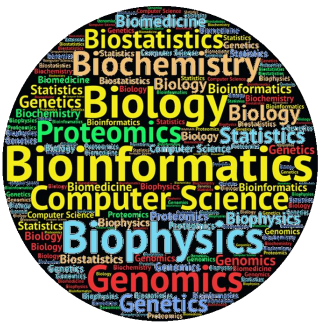
- Familiarizar a los estudiantes en temas de la bioinformática (con sus herramientas, algoritmos y fundamentos)
- 2. Orientar en los usos y aplicaciones de la Bioinformática
- 3. Introducir al sistema operativo LINUX
- 4. Aprender a trabajar por Línea de comandos
- 5. Aprender a utilizar herramientas Bioinformáticas

¿Que es la Bioinformática?

La **bioinformática** es el desarrollo, diseño y la aplicación de tecnologías computacionales a la gestión, administración y análisis de datos biológicos.

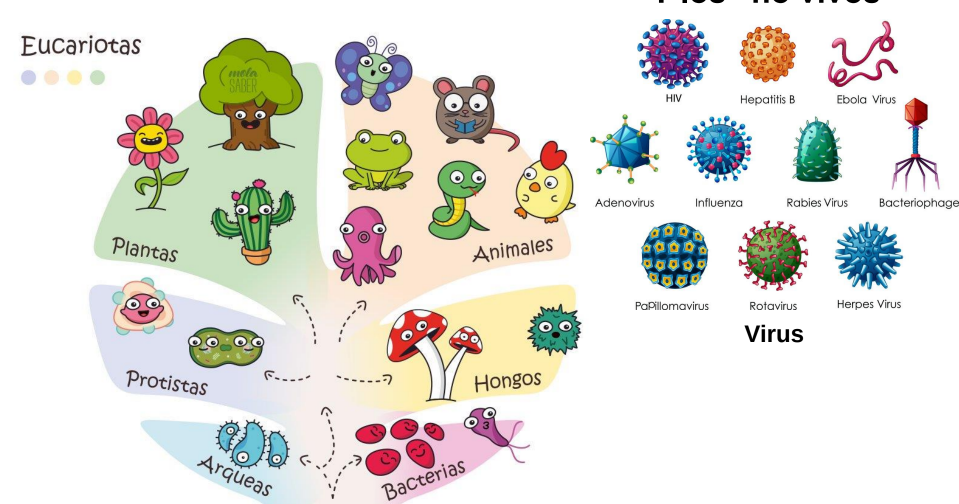
- Biología molecular computacional
- Biología de sistemas computacionales
- Biología Computacional





Reinos de la vida

Y los "no vivos"

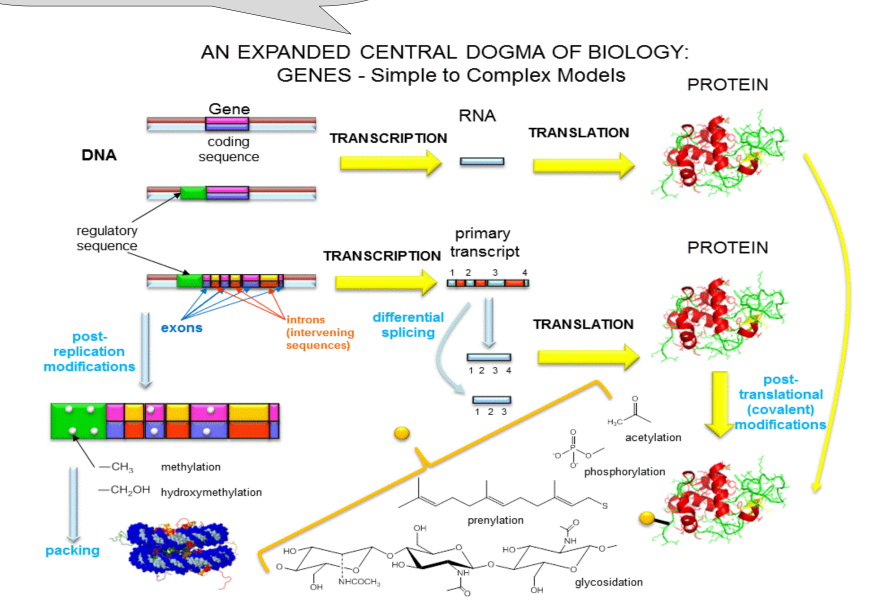






La Bioinformática estudia a todos ellos

Si hay algo que aprendimos en estos ultimos diez años es que tan complejo se volvio todo....



Biology timeline

1962 1953 1955 1970 1972 1975 1977 1977 1980 • Watson & Crick • The sequence of the • Margaret Dayhoff • The details of the • The first recombinant • Two-dimensional •The phi X 174 (or •The full description of •The first complete proposed the double first protein to be (remember that Needleman-Wunsch DNA molecule is electrophoresis, ФХ174) the Brookhaven PDB gene sequence for an helix model for DNA analysed, bovine name!) is renowned algorithm for created by Paul Berg where separation of bacteriophage was (http://www.pdb.bnl. organism (FX174) is based x-ray data insulin, is announed as the first ever sequence comparison and his group. SDS the first genome to gov) is published published. The gene proteins on are published. obtained by Franklin by F.Sanger. bioinformatician. polyacrylamide gel is be sequenced by F. • Allan Maxam and consists of 5,386 base & Wilkins. Recognizing the combined with Sanger. pairs which code nine Gilbert Walter potential for separation according proteins (Harvard) and computational to isoelectric points. Frederick Sanger methods to solve is announced by (U.K. Medical biomedical problems, P.H.O'Farrel. Research Council), in 1962 she released report methods for the first de novo sequencing DNA. sequence assembler in collaboration with Robert Ledley6. The assembler looked at primary protein structure and was in the written programming language Fortran on

University. His thesis Transmission Control

Protocol (TCP).

describes Ethernet.

punch-cards.

Informatics timeline

1974 1969 1973 1975 1976 1978 ARPANET is •The Brookhaven • Vint Cerf and Robert • Microsoft Corporation • The Unix-To-Unix • The Usenet created by linking Protein DataBank is Khan develop the is founded by Bill Copy Protocol (UUCP) connection computers at announeced concept of connecting Gates and Paul Allen is developed at Bell established between Standford and UCLA. (Acta.Cryst.B,1973,29: networks Labs. Duke and the 1764). Robert computers into an University of North Metcalfe receives his "internet" and Carolina at Chapel Hill Ph.D from Harvard by Tom Truscott, Jim develop the

Ellis

Bellovin.

and

Steve

Bioinformatics timeline

The

1990 **BLAST**

1981 Smith-Waterman algorithm for sequence

alignment is published. • IBM introduces its Personal Computer to the market.

The

1986

• Following the Santa Fe conference. DOE OHER announces Human Genome Initiative. - The SWISS-PROT database is created.

1988

- National Center Biotechnology Information (NCBI) created at NIH/NLM
- EMBnet network for database distribution

program (Altschul.et.al.) implemented. Molecular applications group founded in California by Michael Levitt and Chris Lee. The company's products address sequence analysis, database and data management, searching, publication graphics, clone

1994

 Netscape Communications Corporation founded and releases Naviagator, the commerical version of NCSA's Mozilla. Gene Logic is formed in Maryland. The PRINTS database of protein motifs is published by Attwood and Beck. Oxford Molecular Group acquires IntelliGenetics.

1996

The genome Saccharomyces cerevisiae (baker's yeadt, 12.1 Mb) is sequenced. The prosite database is reported by Bairoch, et.al. Affymetrix produces the commerical DNA chips.















construction, mapping and

primer design.









1982

• Genetics Computer Group (GCG) created as a part of the University of Wisconsin of Wisconsin Biotechnology Center. The company's primary product is The Wisconsin Suite molecular biology tools.

- GenBank Release 3 made public
- Phage lambda genome sequenced

1987

 Perl (Practical Extraction Report Language) released by Larry Wall

1989

•The genetics Computer Group (GCG) becomes a privatae company. Primary products: Anaconds, Asp, Cameleon and others (molecular modeling, drug design, protein design).

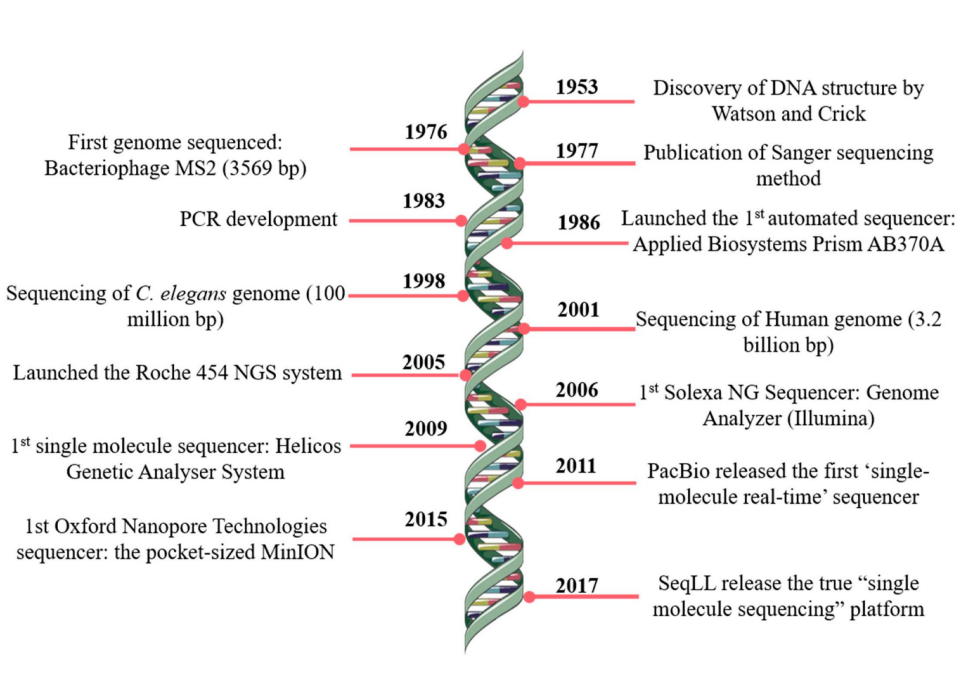
1991

•The research institute in Geneva (CERN) announces the creation of the protocols which make -up the World Wide Web.

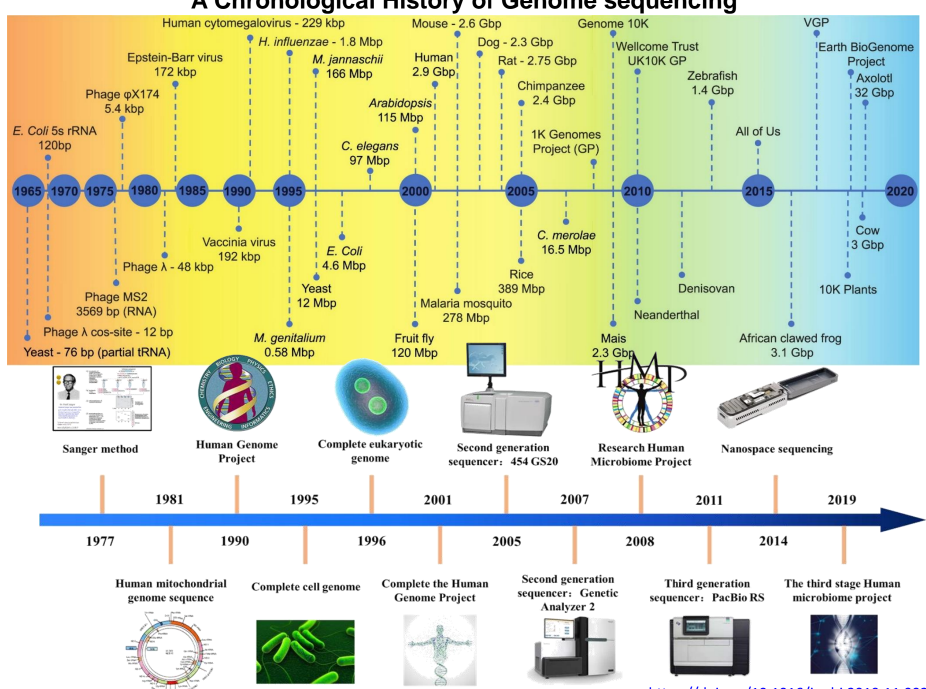
1995

• The Haemophilus influenzea genome (1.8) is sequenced. The Mycoplasma genitalium genome is sequenced.

Milestones in History of Bioinformatics



A Chronological History of Genome sequencing



https://doi.org/10.1016/j.csbj.2019.11.002

Applications of Bioinformatics

- With a large number of prokaryotic and eukaryotic genomes completely sequenced and more forthcoming, access to the genomic information and synthesizing it for the discovery of new knowledge have become central themes of modern biological research.
- Mining the genomic information requires the use of sophisticated computational tools.
 - the careful storage, organization and indexing of information in order to tackle the new challenges in the genomic era.
- Information science has been applied to biology to produce a field is called bioinformatics.
 - It is concerned with the state of- the-art computational tools available to solve biological research problems.
 - The term bioinformatics was coined by Paulien Hogeweg and Ben Hesper to describe "the study of informatic processes in biotic systems" and it found early use when the first biological sequence data began to be shared.
 - Bioinformatics is an interdisciplinary field that develops methods and software tools for understanding biological data.
 - As an interdisciplinary field of science, bioinformatics combines biology, computer science, information engineering, mathematics and statistics to analyze and interpret biological data.
- The key areas of bioinformatics include biological databases, sequence alignment, gene and promoter prediction, molecular phylogenetics, structural bioinformatics, genomics, and proteomics.

Applications of Bioinformatics

- Bioinformatics has become essential for **basic genomic and molecular biology research**, **biotechnology and biomedical sciences**. The main uses of bioinformatics include:
 - Bioinformatics plays a vital role in the areas of structural genomics, functional genomics, and nutritional genomics.
 - It covers emerging scientific research and the exploration of proteomes from the overall level of intracellular protein composition (protein profiles), protein structure, protein protein interaction, and unique activity patterns (e.g. post-translational modifications).
 - Bioinformatics is used for transcriptome analysis where mRNA expression levels can be determined.
 - Bioinformatics is used to identify and structurally modify a natural product, to design a compound with the desired properties and to assess its therapeutic effects, theoretically.
 - Cheminformatics analysis includes analyses such as similarity searching, clustering, QSAR modeling, virtual screening, etc.
 - Bioinformatics is playing an increasingly important role in almost all aspects of drug discovery and drug development.
 - Bioinformatics tools are very effective in prediction, analysis and interpretation of clinical and preclinical findings.

Medical

Molecular medicine

- The human genome will have profound effects on the fields of biomedical research and clinical medicine.
- The completion of the human genome and the use of bioinformatic tools means that we can search for the genes directly associated with different diseases and begin to understand the molecular basis of these diseases more clearly.
- This new knowledge of the molecular mechanisms of disease will enable better treatments, cures and even preventative tests to be developed.

Personalised medicine

- Clinical medicine will become more personalised with the development of the field of pharmacogenomics.
- This is the study of how an individual's genetic inheritence affects the body's response to drugs.
- Today, doctors have to use trial and error to find the best drug to treat a particular patient as those with the same clinical symptoms can show a wide range of responses to the same treatment.
- In the future, doctors will be able to analyse a patient's genetic profile and prescribe the best available drug therapy and dosage from the beginning.

Preventative medicine

• With the specific details of the genetic mechanisms of diseases being unravelled, the development of diagnostic tests to measure a persons susceptibility to different diseases may become a distinct reality.

Gene therapy

- In the not too distant future with the use of bioinformatics tool, the potential for using genes themselves to treat disease may become a reality.
- Gene therapy is the approach used to treat, cure or even prevent disease by changing the expression of a person's genes.

Drug development

- At present all drugs on the market target only about 500 proteins.
- With an improved understanding of disease mechanisms and using computational tools to identify and validate new drug targets, more specific medicines that act on the cause, not merely the symptoms, of the disease can be developed.
- These highly specific drugs promise to have fewer side effects than many of today's medicines.

Antibiotic resistance

- Scientists have been examining the genome of *Enterococcus faecalis*-a leading cause of bacterial infection among hospital patients.
- They have discovered a virulence region made up of a number of antibiotic-resistant genes that may contribute to the bacterium's transformation from a harmless gut bacteria to a menacing invader.
- The discovery of the region, known as a pathogenicity island, could provide useful markers for detecting pathogenic strains and help to establish controls to prevent the spread of infection in wards.

Biotechnology

- The archaeon *Archaeoglobus fulgidus* and the bacterium *Thermotoga maritima* have potential for practical applications in industry and government-funded environmental remediation.
- These microorganisms thrive in water temperatures above the boiling point and therefore may provide the DOE, the Department of Defence, and private companies with heat-stable enzymes suitable for use in industrial processes
- Other industrially useful microbes include, *Corynebacterium glutamicum* which is of high industrial interest as a research object because it is used by the chemical industry for the biotechnological production of the amino acid lysine.
- The substance is employed as a source of protein in animal nutrition.
- Biotechnologically produced lysine is added to feed concentrates as a source of protein, and is an alternative to soybeans or meat and bonemeal.
- Lactococcus lactis is one of the most important micro-organisms involved in the dairy industry.
- Researchers anticipate that understanding the physiology and genetic make-up of this bacterium will prove invaluable for food manufacturers as well as the pharmaceutical industry, which is exploring the capacity of *lactis* to serve as a vehicle for delivering drugs.

The reality of bioweapon creation

- Scientists have recently built the virus poliomyelitis using entirely artificial means.
- They did this using genomic data available on the Internet and materials from a mail-order chemical supply.
- The research was financed by the US Department of Defence as part of a biowarfare response program to prove to the world the reality of bioweapons.
- The researchers also hope their work will discourage officials from ever relaxing programs of immunisation.
- This project has been met with very mixed feelings.

Crop improvement

- Comparative genetics of the plant genomes has shown that the organisation of their genes has remained more conserved over evolutionary time than was previously believed.
- These findings suggest that information obtained from the model crop systems can be used to suggest improvements to other food crops.
- At present the complete genomes of Arabidopsis thaliana (water cress) and Oryza sativa (rice) are available.

Insect resistance

- Genes from *Bacillus thuringiensis* that can control a number of serious pests have been successfully transferred to cotton, maize and potatoes.
- This new ability of the plants to resist insect attack means that the amount of insecticides being used can be reduced and hence the nutritional quality of the crops is increased.

Microbial genome applications

- The arrival of the complete genome sequences and their potential to provide a greater insight into the microbial world and its capacities could have broad and far reaching implications for environment, health, energy and industrial applications.
- For these reasons, in 1994, the US Department of Energy (DOE) initiated the MGP (Microbial Genome Project) to sequence genomes of bacteria useful in energy production, environmental cleanup, industrial processing and toxic waste reduction.
- By studying the genetic material of these organisms, scientists can begin to understand these microbes at a very fundamental level and isolate the genes that give them their unique abilities to survive under extreme conditions

Waste cleanup

- *Deinococcus radiodurans* is known as the world's toughest bacteria and it is the most radiation resistant organism known.
- Scientists are interested in this organism because of its potential usefulness in cleaning up waste sites that contain radiation and toxic chemicals.

Development of Drought resistance varieties

- Progress has been made in developing cereal varieties that have a greater tolerance for soil alkalinity, free aluminium and iron toxicities.
- These varieties will allow agriculture to succeed in poorer soil areas, thus adding more land to the global production base.
- Research is also in progress to produce crop varieties capable of tolerating reduced water conditions.

Climate change Studies

- Increasing levels of carbon dioxide emission, mainly through the expanding use of fossil fuels for energy, are thought to contribute to global climate change.
- Recently, the DOE (Department of Energy, USA) launched a program to decrease atmospheric carbon dioxide levels.
- One method of doing so is to study the genomes of microbes that use carbon dioxide as their sole carbon source.

Alternative energy sources

• Scientists are studying the genome of the microbe *Chlorobium tepidum* which has an unusual capacity for generating energy from light

Evolutionary studies

• The sequencing of genomes from all three domains of life, eukaryota, bacteria and archaea means that evolutionary studies can be performed in a quest to determine the tree of life and the last universal common ancestor.

Comparative Studies

- Analysing and comparing the genetic material of different species is an important method for studying the functions of genes, the mechanisms of inherited diseases and species evolution.
- Bioinformatics tools can be used to make comparisons between the numbers, locations and biochemical functions of genes in different organisms.

Veterinary Science

Sequencing projects of many farm animals including cows, pigs and sheep are now well under way
in the hope that a better understanding of the biology of these organisms will have huge impacts
for improving the production and health of livestock and ultimately have benefits for human
nutrition.

Biological diversity

• The study of different organism and sampling of the ecosystem through the application of new technologies

Contenido de la materia - Clases

Introducción a
Linux – Comandos
básicos y
herramientas
comunes en
Bioinformática

Estructuras de Bases de Datos (DBs) y DBs biológicas communes en Bioinformática

Algoritmos para la comparacion de secuencias

Busqueda de Patrones en secuencias

















Analisis de secuencias:

- Proteinas
- Genes
- Genomas

Comparación de secuencias

Tipos de alineamientos de secuencias y usos

Estructuras de Proteínas y métodos de analisis de estructuras proteicas

Tecnologías de secuenciación por NGS

Genómica

Identificción y estudio de genes

Transcriptómica

















Tipos de datos de NGS y análisis

Estudio de secuencias genómicas

Genómica comparativa

Introdicción a la inteligencia artificial

Introducción a la Bioinformática en Linux

Hardware: Componentes físicos

Generalidades. Definiciones básicas. Hardware

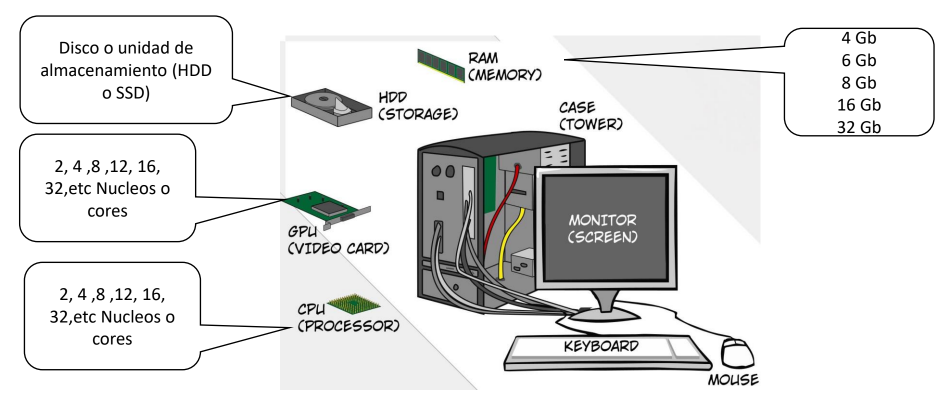
El significado del término **hardware** no es fácil de expresar en español con una sola palabra; literalmente se debe entender como "conjunto de útiles duros"; en el contexto que nos ocupa, el hardware de un computador

Hardware es el conjunto de dispositivos físicos que lo componen,

Software, designa los programas que puede ejecutar el computador.

En cierto modo, el **hardware** es comparable al cerebro o, más generalmente, al cuerpo físico del computador mientras que el **software** será lo equivalente a las ideas que pueblan el cerebro.

Componentes de una computadora



La **memoria de acceso aleatorio** (*Random Access Memory*, **RAM**) se utiliza como memoria de trabajo de <u>computadoras</u> y otros dispositivos para el <u>sistema operativo</u>, los <u>programas</u> y la mayor parte del <u>software</u>. En la RAM se cargan todas las instrucciones que ejecuta la <u>unidad central de procesamiento</u> (procesador) y otras unidades del computador, además de contener los datos que manipulan los distintos programas.

La unidad central de procesamiento o unidad de procesamiento central (conocida por las siglas CPU, del <u>inglés</u>: Central Processing Unit), es el <u>hardware</u> dentro de un <u>ordenador</u> u otros dispositivos programables, que interpreta las <u>instrucciones</u> de un <u>programa informático</u> mediante la realización de las operaciones básicas aritméticas, lógicas y de entrada/salida del sistema.

La <u>unidad de almacenamiento</u> es el <u>lugar</u> donde se guarda la información de manera permanente

What is the Memory Capacity of a Human Brain?

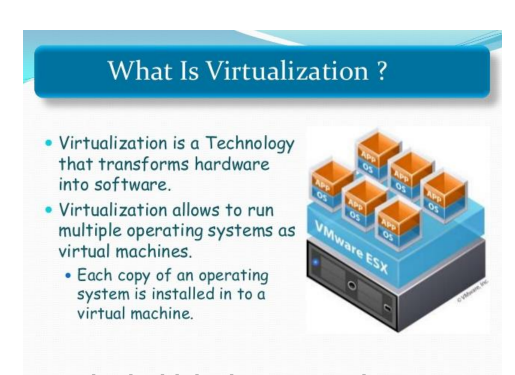


- The human brain's memory capacity in the average adult can store **trillions of bytes** of information.
- In a Stanford Study, it was reported that the cerebral cortex alone has 125 trillion synapses.
- In another study, it was reported that **1 synapse can store 4.7 bits of information**. Neurons are the cells which processes and transmits messages within the brain, and synapses are the bridges between neurons which carry the transmitted messages.
- Running the numbers 125 trillion synapses 4.7 bits/synapse, and about 1 trillion bytes equaling 1 TB (Terabyte).
- This storage capacity is an amount over **74 Terabytes** (just in the cerebral cortex alone)
- according to a 2010 article in Scientific American, the memory capacity of the human brain was reported to have the equivalent of **2.5 petabytes of memory capacity.**

As a number, a "petabyte" means 1024 terabytes or a million gigabytes, so the average adult human brain has the ability to store the equivalent of 2.5 million gigabytes digital memory.

Virtualización y máquinas virtuales

Virtualization refers to the practice of breaking down the physical infrastructure of computing and networking resources into smaller, reusable and more flexible units.



Virtual Machines

Virtual Machines

Virtual Machines

- Method which aims to use the resources effectively
- Isolation of computing resources from users.

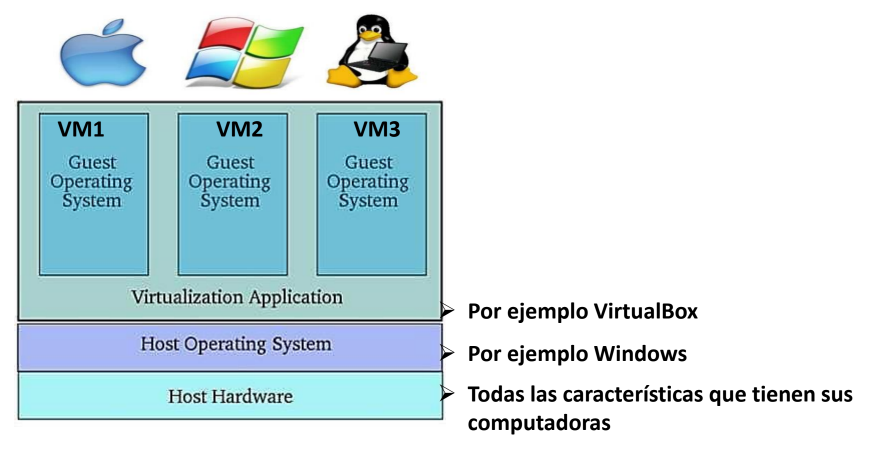
Physical Host Server

Virtualización y máquinas virtuales

- These virtual machines share the physical resource between each other.
- Each virtual machine has own operating system.
- Hardware sharing is set by an interface (Hardware Emulation):
 - Full emulation of complete hardware
 - Supports different hardware components on a
 - different hardware infrastructure
 - Enables each virtual machine to work with its own processor,
 RAM-everything physical computer contains.

OS Virtualization

Guest OS operates in virtual machines within the virtualization application which runs on top of the host OS in the same way as any other application.



Virtualizadores

VirtualBox.org!

VirtualBox is a powerful x86 and AMD64/Intel64 virtualization product for enterprise as well as home use. Not only is VirtualBox an extremely feature rich, high performance product for enterprise customers, it is also the only professional solution that is freely available as Open Source Software under the terms of the GNU General Public License (GPL) version 2. See "About VirtualBox" for an introduction.

Presently, VirtualBox runs on Windows, Linux, Macintosh, and Solaris hosts and supports a large number of guest operating systems including but not limited to Windows (NT 4.0, 2000, XP, Server 2003, Vista, Windows 7, Windows 8, Windows 10), DOS/Windows 3.x, Linux (2.4, 2.6, 3.x and 4.x), Solaris and OpenSolaris, OS/2, and OpenBSD.

VirtualBox is being actively developed with frequent releases and has an ever growing list of features, supported guest operating systems and platforms it runs on. VirtualBox is a community effort backed by a dedicated company: everyone is encouraged to contribute while Oracle ensures the product always meets professional quality criteria.

VMWARE WORKSTATION PLAYER™ 15.5

vmware

VMware Workstation Player

VMware Workstation Player is an ideal utility for running a single virtual machine on a Windows or Linux PC. Organizations use Workstation Player to deliver managed corporate desktops, while students and educators use it for learning and training.

The free version is available for non-commercial, personal and home use. We also encourage students and non-profit organizations to benefit from this offering.

Commercial organizations require commercial licenses to use Workstation Player.

Need a more advanced virtualization solution? Check out Workstation Pro.

¿Como se utiliza la virtualización en bioinformática?

Own computer



S.O Linux

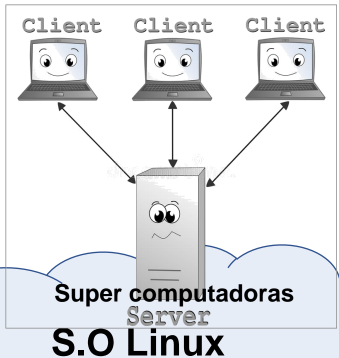
No virtualizado: Utiliza Linux como

sistema operativo principal

Virtualizado: Utiliza Linux como guest en otro sistema operativo

(Limitacion de recursos)

Remote conection



No virtualizado: Utiliza Linux como sistema

operativo principal

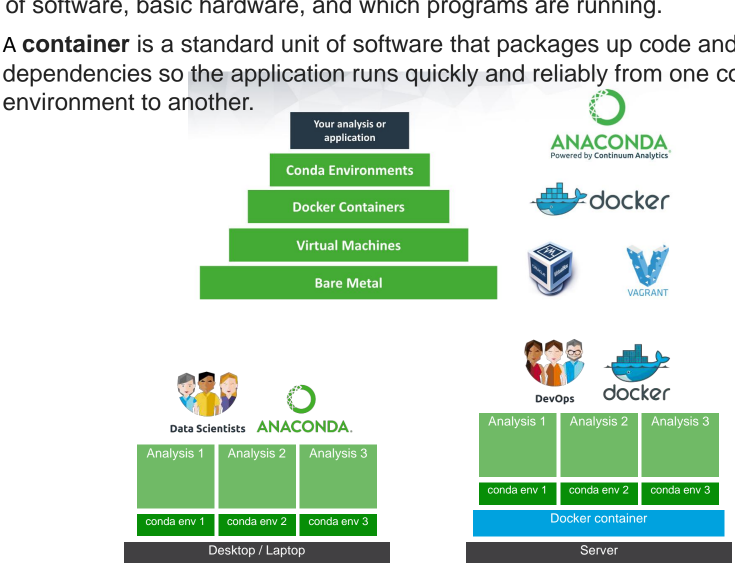
Virtualizado: Utiliza Linux como guest en otro

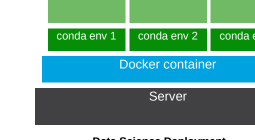
sistema operativo.

Environments and containers

The term **environment** refers to the state of a **computer**, determined by a combination of software, basic hardware, and which programs are running.

A **container** is a standard unit of software that packages up code and all its dependencies so the application runs quickly and reliably from one computing

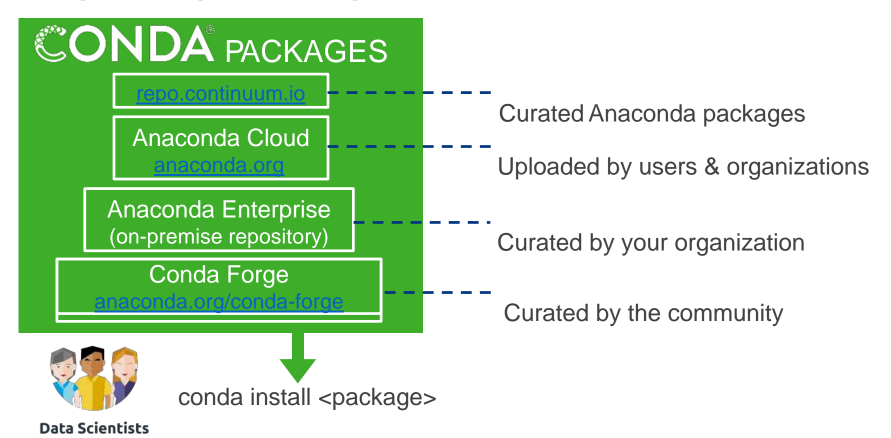




Data Science Development

Data Science Deployment

Creating & Curating conda Packages for Data Scientists





Enforcer of Safety and Correctness

pre-compiled packages only

will never require a compiler in production, or unexpectedly invoke one on you

environment integrity

disk-mutating operations are wrapped in a transaction, and rolled back in the event of errors

environment correctness

conda enforces compatibility of packages within environments

Channels for User Empowerment

- the channel is a component of a package's identity first-class citizen in package specifications
- easy package building
 with conda-build, a dedicated tool with engaged and dedicated code
 contributors
- channels enable community
 independent contributors, independent packaging communities,
 and corporations



Conda is a Community































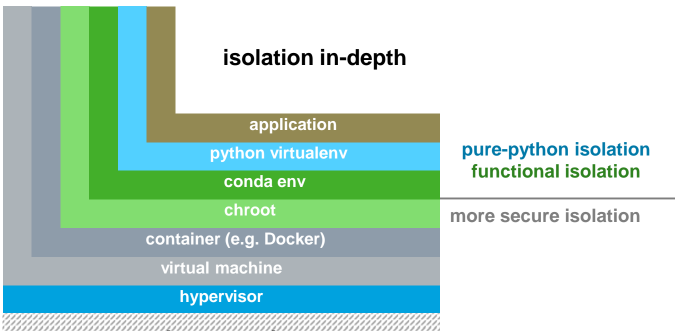






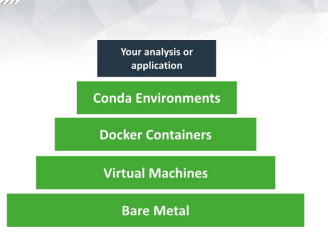
Resumamos por capas

Layers of Process Isolation



bare metal







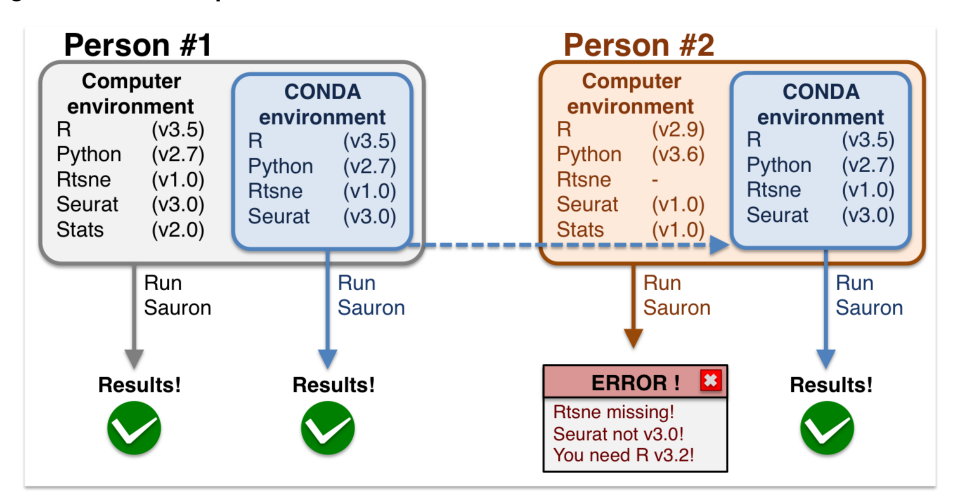






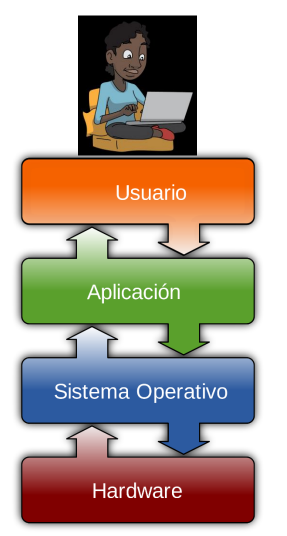
¿Por que usamos un enviroment?

El sistema de *enviroments* trae equilibrio al conjunto de programas para que no haya diferencias o errores en las dependencias de modo que se pueda llegar a un mismo resultado usando los mismos paquetes, las mismas librerías y las mismas versiones de softwares haciéndolo independiente del hardware y del sistema operativo garantizando la reproducibilidad de los resultados



¿Qué es un sistema operativo (OS)?

sistema operativo Conjunto de órdenes y programas que controlan los procesos básicos de una computadora y permiten el funcionamiento de otros programas y la interacción con el usuario.



Dentro de las tareas que realiza el sistema operativo, en particular, se ocupa de **gestionar la memoria de nuestro sistema y la carga de los diferentes programas**, para ello cada programa tiene una prioridad o jerarquía y en función de la misma contará con los recursos de nuestro sistema por más tiempo que un programa de menor prioridad. **El sistema operativo se ocupa también de correr procesos**.

SISTEMA OPERATIVO

De primer plano: requieren de la interacción del usuario, es el caso de un navegador web, un editor de texto, un programa de diseño de imágenes.

De segundo plano: son aquellos programas que no requieren del usuario y habitualmente no poseen una interfaz gráfica o "pantalla". Un ejemplo de este tipo de procesos es el anti-virus.

Es libre

La principal ventaja que tienen muchos sistemas operativos basados en Linux es que es software libre y de código abierto. Eso significa, en la práctica, que todos pueden acceder al código fuente del sistema operativo, revisarlo y contribuir a su desarrollo.

Es abierto

Podemos tocar, ajustar y tunear Linux hasta niveles insospechados. Desde elegir el entorno de escritorio que queremos utilizar

Es (generalmente) más seguro

Hace unos años sucedió que las Fuerzas Armadas de los Estados Unidos sufrieron el ataque de un virus informático que inmovilizó algunos sistemas tan críticos como el que controla algunos de sus aviones no tripulados, como el Predator. ¿A qué sistema operativo migraron? A uno basado en Linux.

Está en todos lados

Aunque no lo veamos, Linux realmente está en todas partes. Hay sistemas operativos que usan el núcleo de Linux aunque no lo parezca. Ej. plataformas que usan Linux:

- Android
- Chrome OS
- •webOS (en Palm, en algunos dispositivos de HP, y en televisores de LG)
- •Tizen
- Steam OS

Puede revitalizar tu ordenador antiguo

El núcleo de Linux es extremadamente ligero y, junto con un entorno de escritorio sin demasiada floritura, **es capaz de funcionar en ordenadores que tengan más de diez años**.

¿Por qué Linux es mejor para programadores y desarrolladores?

1.Código abierto

Linux es un sistema operativo de código abierto, lo que significa que está abierto al público. **Un programador puede ver y editar o contribuir al código fuente que se usó para crearlo.** Pueden crear su propia versión del sistema operativo que puede ayudarlos con áreas especializadas o estratégicas. Otros países también están desarrollando sus propios sistemas operativos basados en el código fuente.

2.Sin restricciones

Linux no tiene restricciones. No es necesario que aguardes interminablemente una actualización de funciones o un parche de seguridad y obstáculos para actualizar la licencia de usuario. Un sistema Linux es muy estable ymenos propenso a malware y virus. Se puede usar en instituciones educativas, en casa. En cualquier sitio.

3. Estructura configurable

Linux está estructurado como capas (Kernel, hardware, IO y UI), que son extremadamente configurables. Puedes cambiar algo si no te gusta la forma en que está funcionando. Linux no requiere una interfaz gráfica de usuario para interactuar, es decir, solo con la línea de comando (Shell). Un sistema Linux se puede ajustar con precisión para aprovechar las posibilidades máximas del hardware.

Como código de fuente abierta, los usuarios tienen derecho a ver y modificar el código fuente e incluso crear el suyo propio.

4. Menos requisito de hardware

Linux es muy eficiente en términos de los recursos del sistema. Dado que Linux se puede personalizar, la instalación para los usuarios y los requisitos específicos de hardware son fáciles. El procedimiento de instalación flexible permite a los usuarios elegir lo que quieran instalar. Esto incluso permite a los programadores instalar Linux en hardware antiguo y permite el uso óptimo de todos los recursos de hardware disponibles.

5. Disponibilidad de las herramientas de programación

Linux es gratis, es por eso que todo el software básico (que es necesario para un usuario típico o incluso un usuario avanzado) está disponible. Hay muchos programas educativos disponibles bajo Linux. Las contrapartes de software profesional para autoedición, edición de fotos, edición de audio y edición de videotambién están disponibles.

¿Por qué Linux es mejor para programadores y desarrolladores?

6. Grandes comunidades de soporte

Dado que Linux ya existía desde hace más de 26 años, a lo largo de los años había creado un fuerte apoyo comunitario. Los programadores pueden encontrar soporte fácilmente a través de Internet. Los foros y otros sitios web entusiastas de Linux pueden ayudarte de inmediato cuando tengas alguna pregunta en mente. Sin duda, nunca estarás solo con este sistema operativo.

7. Fácil de instalar

Como programador, instalar el sistema operativo Linux nunca debería ser un problema. Presionar las teclas necesarias durante el arranque y navegar por las indicaciones en pantalla será fácil porque Linux no es tan diferente de instalar un programa desde otros sistemas operativos.

8.Demanda

La demanda de Linux es excelente. Dado que es de código abierto y gratuito, cada vez más programadores desean crear su propia versión de Linux que pueda ser exitosa en el futuro y pueda ayudar a particulares o incluso al gobierno.

9. Actualizaciones fáciles

Las actualizaciones en Linux son muy fáciles. Como todo el tipo de Linux tiene su propio repositorio central de software. Actualizaciones regulares que están disponibles e incluso el sistema se puede actualizar sin reiniciarlo. Los usuarios pueden incluso automatizar el proceso de actualización. A diferencia de Windows, la actualización consume menos almacenamiento y datos de Internet.

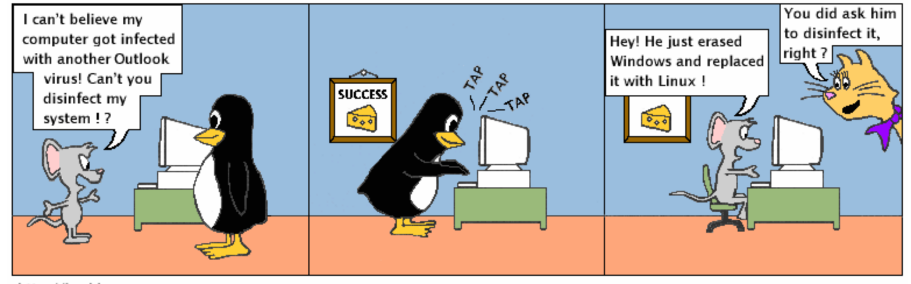
10.Personalización

Linux incluye diferentes tipos de opciones para el software. Esto significa que las aplicaciones como losprocesadores de texto y los navegadores web se pueden cambiar según tu elección. Los usuarios de Linux también pueden elegir pantallas de sistema, gráficos y otros componentes de interfaz de usuario que mejor se adapten a su imagen. La idea es que, con Linux, los programadores pueden usar varios programas pequeños pero brillantes que se pueden combinar para escribir programas realmente poderosos y utilidades para que los usen.



Hackles

By Drake Emko & Jen Brodzik



http://hackles.org

GNU/Linux

GNU/Linux es la denominación técnica y generalizada que reciben una serie de sistemas operativos de tipo Unix, que también son multiplataforma, multiusuario y multitarea, pero libres. 1 Estos sistemas operativos están formados mediante la combinación de varios proyectos, entre los cuales destaca el entorno GNU, encabezado por el programador estadounidense Richard Stallman junto a la Free Software Foundation, una fundación cuyo propósito es difundir el software libre, así como también el núcleo de sistema operativo conocido como «Linux», encabezado por el programador finlandés Linus Torvalds.²

https://histinf.blogs.upv.es/2011/12/23/historia-de-linux/

















Server oriented

RedHat, CentOS, openSuse, Ubuntu Server, Debian, ...













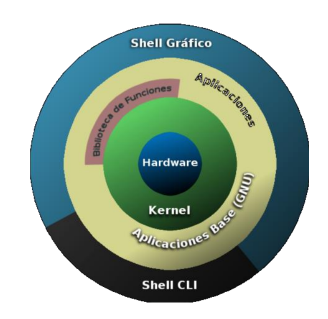




Home oriented

Fedora, Linux Mint, openSuse, Ubuntu, Gentoo, Android,

UNIX OS – Algunos conceptos



What?

 Collection of software that manages computer hardware resources and provides common services for programs

Kernel

- An OS kernel manages computer resources (e.g. CPU, RAM, internal/external devices) and allows programs to use
- these resources

Shell

• = Command Line Interpreter (CLI) is a text-only interface between the user and the kernel. Its function is to execute commands from the terminal window

Programs

External programs can be installed. The OS comes with many built-in utilities

EVENTS LEADING TO CREATION OF LINUX

- 1969: Ken Thompson, de AT&T Bell Laboratories, desarrolló el sistema operativo Unix, adaptándolo a las necesidades de un entorno de investigación 1975: Berkeley lanzó su propia versión de Unix (BSD). Esta versión de Unix se convirtió en la
- principal competidora de la versión de los laboratorios Bell de ATT&T

 1980-1990: se fragmentó BSD, fue superado por Linux y dio origen a Mac OS X
- 1980-1990: se fragmentó BSD, fue superado por Linux y dio origen a Mac OS X
 1983: Richard Stallman started the GNU project with the goal of creating a free UNIX-like operating system. As part of this work, he wrote the GNU General Public License (GPL)
 1991: Linus Torvalds announced Linux Kernel Linus anuncio la primera versión oficial de Linux (versión 0.02). Con esta versión Linus pudo ejecutar Bash (GNU Bourne Again Shell) y gcc (El

https://histinf.blogs.upv.es/2011/12/23/historia-de-linux/

https://itsfoss.com/25-years-linux/

- compilador GNU de C)

 1992: Linux became Open Source
- 1994: Someone registered Linux trademark and it was not Linus Torvalds
- 1995: First Linux Expo
- 1996: Tux gets to be the symbol of Linux 1997: GNOME Project is born
- 1998: KDE 1.0 released
- 2000: Steve Jobs made an offer that Linus refused
- 2002: Red Hat Enterprise Linux released
- 2003: Attempt to install backdoor in Linux kernel 2004: Ubuntu 4.10 released
- 2004: Ubuntu 4.10 released 2005: Linus Torvalds created Git
- 2007: Linux powered netbook arrived (Asus)
- 2008: Android version 1.0 released
- 2009: Google announced Chrome OS (Chromebooks)
- 2010: Red Hat became first billion dollar open source company
- 2011: Microsoft was one of the top 5 contributors to Linux Kernel 2015: Microsoft has its own version of Linux

¿Por qué Linux es mejor para Bioinformática?

- Es libre, de código abierto y gratuito
- Es liviano, estable y permite optimizar recursos
- Funciona en prácticamente todos los computadores
- Permite trabajar con un gran volumen de datos
- Permite concatenar procesos de manera que los programas trabajen en serie (pipelines)
- Tiene gran soporte de la comunidad de programadores y científica
- Permite programa tu propio código en diferentes lenguajes de programación

One definition of bioinformatics is "the use of computers to analyze biological problems." As biological data sets have grown larger and biological problems have become more complex, the requirements for computing power have also grown.

Computers that can provide this power generally use the Unix operating system -

http://bl831.als.lbl.gov/~jamesh/powerpoint/unix_commands.ppt

Slightly more advanced: http://bl831.als.lbl.gov/~jamesh/elves/manual/tricks.html

Basic unix commands that everyone should know

(Even if you have a mac)

What the ~*&?!

```
"tilde" indicates your home directory: /home/you
\sim
*
     "star": wildcard, matches anything
     wildcard, matches any one character
     History substitution, do not use
\delta
     run a job in the background, or redirect errors
     special characters for most crystallography programs
                  back-quote, backslash, etc. special to shell
     underscore, use this instead of spaces!!!
```

Where am I?



Print name of the "current working directory"

This is the default directory/folder where the shell program will look first for programs, files, etc. It is "where you are" in Unix space.

What is a directory?

/home/yourname/whatever

Directories are places you put files. They are represented as words connected by the "/" character. On Windows, they use a "\", just to be different. On Mac, they are called "folders". Whatever you do...

DO NOT PUT SPACES

In directory/file names!

What have we here?

ls

List contents of the current working directory

```
    ls -l
    ls -lrt
    ls -lrt
    ls -lrt /home/yourname/something
```

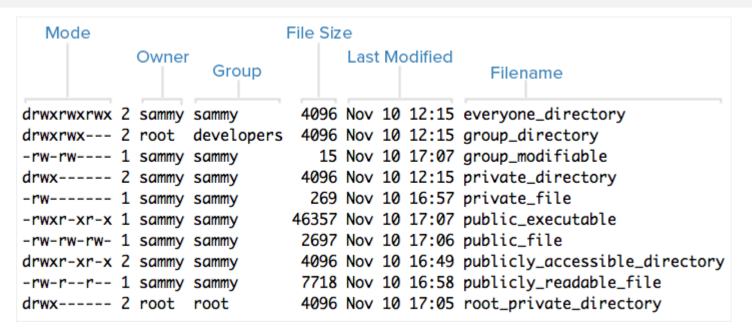
long-list a different directory

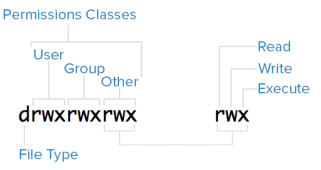
Viewing Ownership and Permissions

In Linux, each and every file is owned by a single user and a single group, and has its own access permissions. Let's look at how to view the ownership and permissions of a file.

The most common way to view the permissions of a file is to use $1\mathsf{s}$ with the long listing option, e.g. $1\mathsf{s}$ -1 myfile . If you want to view the permissions of all of the files in your current directory, run the command without an argument, like this:

Is -I





Permissions Classes

From the diagram, we know that *Mode* column indicates the file type, followed by three triads, or classes, of permissions: user (owner), group, and other. The order of the classes is consistent across all Linux distributions.

Let's look at which users belong to each permissions class:

•User: The owner of a file belongs to this class

•Group: The members of the file's group belong to this class

•Other: Any users that are not part of the *user* or *group* classes belong to this class.

Here is a quick breakdown of the access that the three basic permission types grant a user:

Read

For a normal file, read permission allows a user to view the contents of the file.

For a directory, read permission allows a user to view the names of the file in the directory.

Write

For a normal file, write permission allows a user to modify and delete the file.

For a directory, write permission allows a user to delete the directory, modify its contents (create, delete, and rename files in it), and modify the contents of files that the user has write permissions to.

Execute

For a normal file, execute permission allows a user to execute a file (the user must also have read permission). As such, execute permissions must be set for executable programs and shell scripts before a user can run them.

For a directory, execute permission allows a user to access, or traverse, into (i.e. Cd) and access metadata about files in the directory (the information that is listed in an 1s-1).

Permissions Classes

Examples of Modes (and Permissions)

Now that know how to read the mode of a file, and understand the meaning of each permission, we will present a few examples of common modes, with brief explanations, to bring the concepts together:

- •- rW-----: A file that is only accessible by its owner
- •- rwxr-xr-x: A file that is executable by every user on the system. A "world-executable" file
- •- rw-rw-rw-: A file that is open to modification by every user on the system. A "world-writable" file
- drwxr-xr-x: A directory that every user on the system can read and access
- drwxrwx---: A directory that is modifiable (including its contents) by its owner and group
- drwxr-x---: A directory that is accessible by its group

"Permission denied"!?

chmod

Change the "permission" of a file.

chmod a+r filename.txt

- make it so everyone can read it
- chmod u+rwx filename.txt
 - make it you can read/write/execute it
- chmod -R u+rw /some/random/place
 - make it so you can read/write everything under a directory

Go somewhere else?

cd

Change the current working directory

```
    cd /tmp/yourname/
    go to your temporary directory
    cd - go back to where you just were
    cd - no arguments, go back "home"
    "home" is where your login starts
```

A new beginning...

mkdir

Create a new directory.

```
mkdir ./something- make it

cd ./something - go there

ls - check its is empty
```

How do I get help?

man

Display the manual for a given program

```
man ls - see manual for the "ls" command
man tcsh - learn about the C shell
man bash - learn about that other shell
man man - read the manual for the manual
```

to return to the command prompt, type "q"

Move it!



Move or rename a file. If you think about it, these are the same thing.

```
mv stupidname.txt bettername.txt
    -change name
mv stupidplace/file.txt ../betterplace/file.txt
```

mv stupidname_*.img bettername_*.img

- same name, different directory

Will not work! Never ever do this!

Copy machine

cp

Copy a file. This is just like "mv" except it does not delete the original.

cp stupidname.txt bettername.txt

- change name, keep original

rm stupidname.txt

- now this is the same as "mv"

Destroy! Destroy!



Remove a file forever. There is no "trash" or "undelete" in unix.

```
rm unwanted_file.txt
```

- delete file with that name

```
rm -f /tmp/yourname/*
```

- forcefully remove everything in your directory.

temporary

Will not prompt for confirmation!

less is more

more

Display the contents of a text file, page by page

```
more filename.txt - display contents
less filename.txt - many installs now have a
replacement for "more" called "less" which has nicer search
features.
```

to return to the command prompt, type "q"

After the download...

gunzip

File compression and decompression

gunzip ~/Downloads/whatever.tar.gz

- decompress

gzip ~/Downloads/whatever.tar

- compress, creates file with .gz extension

Where the %\$#& is it?

find

Search through directories, find files

```
find ./ -name 'important*.txt'
```

- look at everything under current working directory with name starting with "important" and ending in ".txt"
- find / -name 'important*.txt'
 - will always find it, but take a very long time!

Did I run out of disk space?

df du

Check how much space is left on disks

- df look at space left on all disks
- df . look at space left in the current working directory
- du -sk . | sort -g
- add up space taken up by all files and subdirectories, list biggest hog last

Why so slow?

ps top

Look for programs that may be eating up CPU or memory.

top

- list processes in order of CPU usage
- jobs **terminal**
- list jobs running in background of current

ps -fHu yourname

- list jobs belonging to your account in order of what spawned what

Die Die Die!

kill

Stop jobs that are running in the background

```
kill %1 - kill job [1], as listed in "jobs"
kill 1234 - kill job listed as 1234 by "ps" or "top"
kill -9 1234 - that was not a suggestion!
kill -9 -g 1234 - seriously kill that job and the program that launched it
```

Linux Cheat sheet by A.Mahouachi 1 File Commands ls [options] file options -a: show hidden files -A: show hidden files except . and .. -d: only show directories -h: human readable size -i: inode info -l: long list format -m: output as csv -n: numeric uid and guid -r: sort in reverse order -S: sort by file size t: sort by modification time tree [options] dir options -d: only directories -f: show full paths -P pattern: only matching pattern -I pattern: except matching pattern -h: print sizes in human readable format -C: use colors -L max: max level depth cp [options] source dest -b: backup dest before overwrite -l: link files instead of copy -P: dont follow sym links

options

-r: recursive -f: force

-i: interactive

u: copy only if source newer than dest

mv [options] source dest options

-b: backup dest before overwrite -f: force

i: interactive

-u: move only if source newer than dest

In [options] file link

options

-s: sym link (hard by default) -f: overwrite link if exists

-b: backup old link before overwrite

rm [options] file

options -f: force

-i: interactive rm - -foo if file name is -foo

chmod [options] mode file(s)

options -R: recursive

symbolic mode

format: [ugoa][[+-=][perms]],... example: u+x,o-wx,g-w

u: owner g: group

o: others a: all +: add mode -: remove mode

first digit: setuid(4), setguid(2)

find path [options] [tests] [actions]

-name "xyz*": name like xyz*

-type d: only directories

-mtime 0: modified < 1 day

-size +100M: size > 100mb

-print: print matching

-delete: rm matching files

-mtime -x: modified < x days</p>

-mtime +x: modified > x days

-perm /o+w: writable by others

-mmin: like -mtime but in minutes

-size -100M: size < 100mb (k for kb, G</p>

!-perm /o+r: not readable by others

-exec cmd '{}' ; : run cmd for every

-exec cmd '{}' + : run cmd at the end of

-exec rm -rf ": rm -rf matching items

-fprint /tmp/result: write matches to

-x".sync*": exclude files with path like

-type f:only files

-mindepth: start from min level in hier-

-maxdepth: end with max level in hier-

-iname "xyz*": like -namé but case in-

second digit: owner perms

third digit: group perms

fourth digit: others perms

s: setuid or setguid

t: sticky bit

numeric mode

example: 755

read: 4

write: 2

options:

archy

tests:

for gb)

actions:

match

search

options

options

-i: ignore case

/tmp/result

-r: recursive

diff [options] files

-w: ignore whitespaces

-q: only show file names

grep [options] pattern files

B: ignore blank lines

execute: 1

format: [0-7]1,4

 R: recurse directories -c: only show matching lines count =: exact mode -exclude=glob : exclude these r: read

w: write -include=glob : only consider these x: execute files and search for dirs cat [options] file(s) X: search for dirs

options

-v: non ascii chars except tab and eol

-n: also show matching line number

-T: show tabs

-t: equivalent to -vT -E: show eol end of line

-P: pattern is a perl regex

-m: stop after m matches

-e: equivalent to -vE -A: equivalent to -vET

-s: remove repeat empty lines

tail [options] file

options

-f: show end of file live -35: show last 35 lines

-q: be quiet head [options] file

options

-35: show first 35 lines

-q: be quiet tac file(s)

print files starting from last line

cut [options] file options

-d char: use char as delimiter -f 1,3,5: print fields 1, 3 and 5

unia (options) input output

options

-c: prefix lines by number of occur-

 -d: only print duplicate lines -u: only print unique lines

sort [options] file options

n: numeric sort

-b: ignore blank lines -f: ignore case

-r: reverse order

tar [options] file

options

-f file: archive file

-c: create -t: list

-x: extract

-C DIR: cd to DIR -z: gzip

-j: bzip2

du [options] file

options:

-c: a grand total -h: human readable

 L: dereference sym links -P: no dereference of sym links

-s: total for each argument

-exclude=pattern -max-depth=N: dont go deeper than N df [options] file

options:

-h: human readable -i: list inodes info

-P: no dereference of sym links

2 Process Commands

ps [options]

options:

-e: all processes -f: full listing -H: show hierarchy -p pid: this process pid

C cmd: this command name cmd

-w: wide output -ww: to show long command lines

-l: long listing, including wchan

-o x,y,z: show columns x y z

-o user,pid,cmd: show columns user, pid command

-N: negation

-u user: processes owned by user -u user -N: processes not owned by user

-sort=x,y: x y are columns in ps output -sort=user: sort by user

-sort=+time: sort by cpu time asc -sort=-time: sort by cpu time desc -sort=size: sort by memory size

-sort=vsize: sort by vm size

top [options]

options

-d x: refresh every x seconds -p pid1 -p pid2: only processes with pid1 pid2 -c : show command lines

interactive commands space: udpdate display

n: change number of displayed proces-

up and down: browse processes k: kill a process

o: change order T: sort by time

A: sort by age P: sort by cpu

M: sort by memory c: display/hide command line m: display/hide memory

t: display/hide cpu

f: manage list of displayed columns up and down: move between columns d: display/hide the selected column q: apply and quit the field mgmt screen

pgrep [options] pattern options

 -1: show pid and process name -a: show pid and full command line

-n: if more than one show newest

 -o: if more than one show oldest u uid : show only processes of uid -c : count results

3 Network & Remote

ssh [options] user@host ["cmd1:cmd2"] options:

-2: force protocol 2

-o StrictHostKeyChecking=no: ignore warnings due to remote host key change

-X: forward X11 display

wget [options] url

options:

-b: run in background

-o file: print wget output in file

-o /dev/null: suppress wget output -q: be quiet

-d: debug

-O file: save response to file

-c: resume file download -S: print server headers

-T N: timeout after N seconds

-user=user: basic http auth user

-password=password: basic http auth

-save-cookies file: save cookies to file -load-cookies file: use file as cookies

-post-data=string -post-file=file

-no-check-certificate: ignore ssl certifi-

curl [options] url

options:

-H header: like -H "Host: st.com"

-u <user:password>: basic http auth -s: be silent

-S: show errors if silent mode

 L: follow new location in case 301 –data "field=value": x-www-form-

urlencoded query -data-binary data: post data as is without encoding

-data-binary @filename: post filename content as is

 X method: use PUT, GET, POST etc. -request method: use PUT, GET, POST

etc. mail [options] to-address

options:

s subject: email with subject

-c address1.address2: cc copy -b address1,address2: bcc copy mail -s 'hello there' 'joe@st.com' < so-

mefile 4 Terminal

Ctrl+C: halt current command Ctrl+Z: pause current command

bg %1: resume paused command in

fg %1: resume paused command in foreground

Ctrl+D: logout

Ctrl+W: remove a word from current

Ctrl+U: remove current line

Ctrl+A: go to beginning of current line Ctrl+E: go to end of current line