BITAH05 -Databanktechnologie

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Overview

The student is able to

- Solve exercises on both normalisation and database creation
- Create a database model
- Recognise several database types
- Use online databases

Course material

- Slides
- Syllabus

Examination

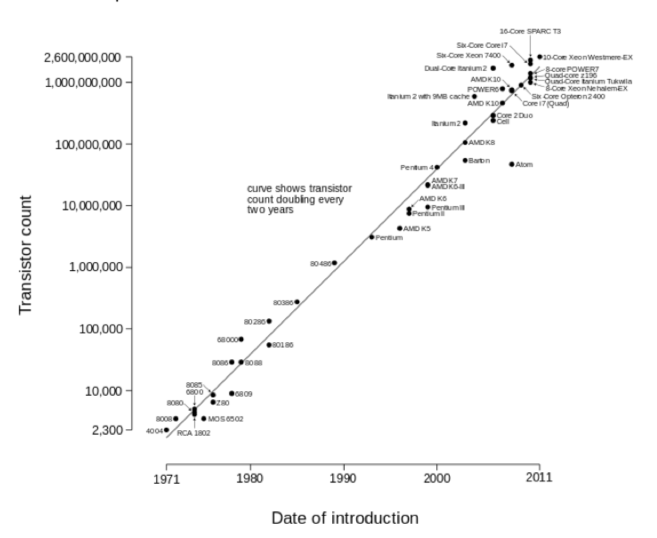
Theoretical and practical part

Requirements

A working internet connection

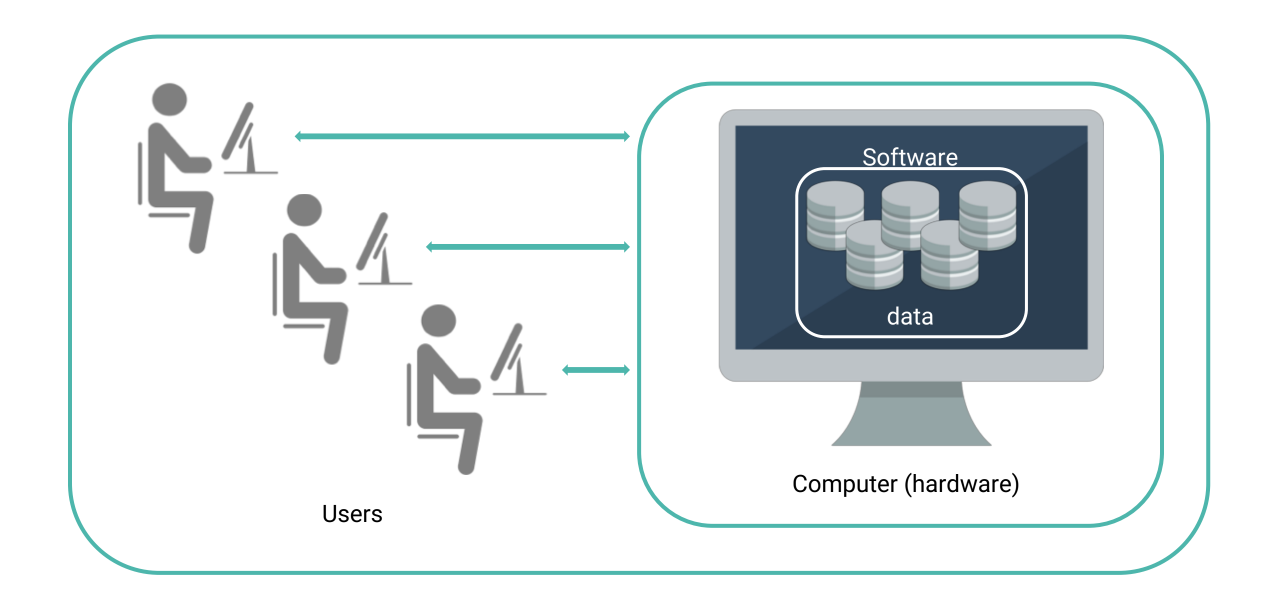
Lecture 1 – Introduction & relational databases

Microprocessor Transistor Counts 1971-2011 & Moore's Law



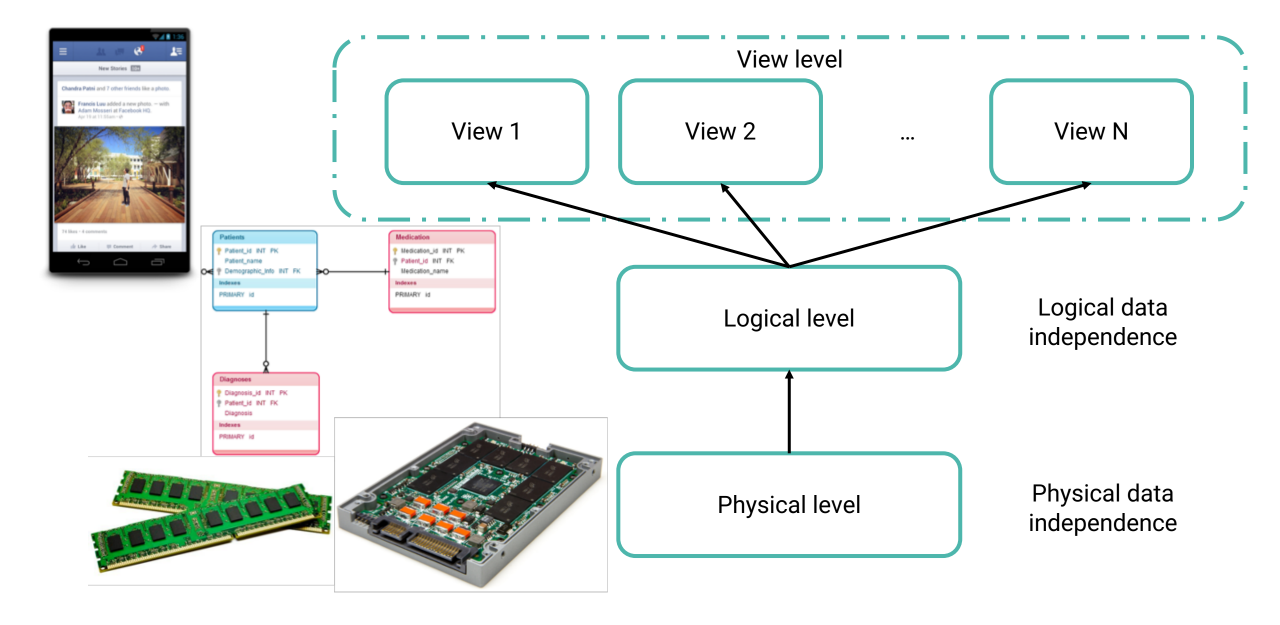
Database

- Mechanism used to store information
- Collection of data (numbers, dates, text, ...)
- Structured storage of data
- Efficient interaction with data
 - CRUD
- Used everywhere!



Database system

- Hardware
 - Processor and internal memory
 - Personal laptop server cluster
- Data
 - Structured storage
 - Little redundancy
 - Shared (single-user vs. multi-user)
- Software
 - Database Management System (DBMS)
 - Data storage
 - Data retrieval
 - Data manipulation
- Users
 - Authentication & authorization
 - Administrator > end-user



A good database system

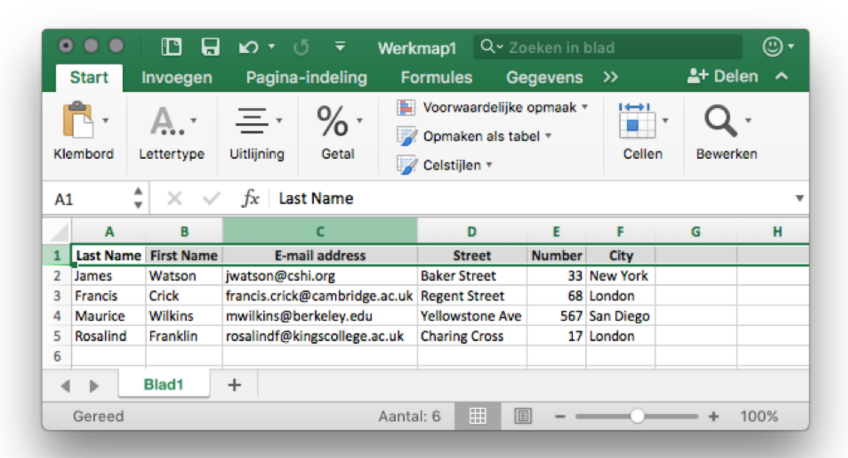
- Sufficient amount of storage
 - Think ahead
- Easily accessible
 - Centralised system
- Secured against non-users
 - Per user rights
- Easily maintainable
 - Add, edit, delete data
- Controlled input
 - All data in same format
- Quick!
 - Queries need to be run within a certain time
- Little redundancy
 - Do not store the same information twice (or more)
 - Application dependable
- Clear structure

Database Management System (DBMS)

- A database management system (DBMS) is a computer software application that interacts with the user, other applications, and the database itself to capture and analyse data
- Shields users from hardware and storage details
- Most important software component, though not the only one (development tools, ...)
- Used for
 - Data storage
 - Data retrieval
 - Data manipulation
 - Authentication & authorization

Relational databases

- Rigid structure
- 2 dimensional tables
 - Columns (fields)
 - Rows (records)



Relational databases

- Model objects (entities) and their relationships
- E.g. a store sells products to customers
 - Entities:
 - Customers

Attributes: name, address, telephone number, ...

Products

Attributes: name, price, ...

- Relationships:
 - Sale

Attributes: quantity, timestamp, ...

Relational Database Management Systems (RDBMS)

- Enforce data integrity
 - Honours constraints on columns
- Enforce referential integrity
 - Honours constraints on relations

! 12 rules of Edgar Codd!

Relational Database Management Systems (RDBMS)

- Commercial products
 - Oracle
 - DB2 (IBM)
 - MS SQL Server (Microsoft)
- Open-source
 - MySQL (Oracle)
 - PostgreSQL
 - SQLite

Relational Database with MySQL

- Most used RDBMS
- Open source
- Free of charge (paying versions exist)
- Wordpress, Twitter, Facebook, ...
- http://www.mysql.com



Installing MySQL on your system

- Linux
 - http://dev.mysql.com/doc/refman/8.0/en/linux-installation.html
- Mac
 - http://dev.mysql.com/doc/refman/8.0/en/osx-installation.html
- Windows
 - http://dev.mysql.com/doc/refman/8.0/en/windows-installation.html

Starting/stopping/restarting MySQL

Linux/Mac

```
$ /etc/init.d/mysqld start service mysqld start service mysqld start
$ /etc/init.d/mysqld stop service mysqld stop service mysql stop
$ /etc/init.d/mysqld restart service mysqld restart service mysqld restart
```

Windows

```
$ C:\> "C:\Program Files\MySQL\MySQL Server 5.7\bin\mysqld"
$ C:\> "C:\Program Files\MySQL\MySQL Server 5.7\bin\mysqladmin" -u root shutdown
```

Check whether or not MySQL is running correctly

Linux/Mac

```
$ service mysqld status
mysql start/running, process 3394
$ ps -ef | grep mysql
mysql 3394 1 0 12:09 ? 00:00:00 /usr/sbin/mysqld
$ netstat -ltpn | grep mysql
tcp 0 0 0.0.0:3306 0.0.0:* LISTEN 3394/mysqld
```

Windows

```
$ C:\> "C:\Program Files\MySQL\MySQL Server 5.7\bin\mysqlshow"
$ C:\> "C:\Program Files\MySQL\MySQL Server 5.7\bin\mysqlshow" -u root mysql
$ C:\> "C:\Program Files\MySQL\MySQL Server 5.7\bin\mysqladmin" version status proc
$ C:\> "C:\Program Files\MySQL\MySQL Server 5.7\bin\mysql" test
```

Or check running services through Control Panel

The MySQL monitor

• To connect or log on to a MySQL database service

```
$ mysql
```

Many options, check the manual

```
$ man mysql
or
$ mysql --help
```

The MySQL monitor

Exercises

Connect to the database and execute the following commands

```
mysql> select current_user;
mysql> show databases;
```

Securing the server

```
$ mysql_secure_installation
```

- Set password for root accounts
- Remove anonymous-user accounts
- Remove remote root login
- Remove test database

Securing the server – extra

- Prevent any external access to the database server
 - Add to global config file (/etc/mysql/my.cnf)

[mysqld]
bind-address = 127.0.0.1

Database users

- Database users and OS users are completely independent from each other
 - No user specified --> OS user is taken
 - Database superadmin --> root@localhost (all rights, including dropping databases)
- Not a good idea to always connect as root!!!

Database user - privileges

- Created user has very limited privileges
- Grant privileges prv on table tbl in database db mysql> GRANT prv ON db.tbl TO user@host;
- Some wild cards
 - All privileges, specify all as prv
 - All databases, specify * as db
 - All tables, specify * as tbl
- The given database and table names do not have to exist (yet)

The options file

- Avoid retyping of password with every connection, create options file
 - .my.cnf
 - Located in home directory
 - Protect from others: mode 600
- Contains *key=value* pairs in [sections]
 - Provided as (invisible) command line parameters

The options file - example

- Put password in options file
 - Command line parameters of mysql

```
$ mysql --password=pwd
```

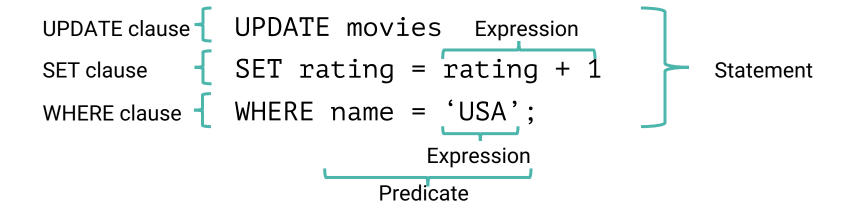
Options file could look like this

```
[client]
password=pwd
user=username
```

SQL: Structured Query Language

- Data Definition Language (DDL) statements: design (create, alter, drop, ...) database CREATE TABLE, DROP DATABASE
- Data Manipulation Language (DML) statements: manage data
 - Create: add new data
 - Read: collect data
 - **U**pdate: alter data
 - **D**elete: remove data SELECT, INSERT, UPDATE, DELETE
- **Data Control Language** (DCL) statements: manage database rights GRANT, REVOKE
- Transaction Control Language (TCL) statements: manage DML tasks (group, undo, ...)

SQL: Structured Query Language



The MySQL monitor (again)

- Several ways to execute SQL statements using the MySQL monitor
 - Interactively

```
$ mysql [database]
mysql> stmt
```

From the command line

```
$ mysql [database] -e 'stmt'
```

From a file or a pipe (stdin)

```
$ mysql [database] < stmt_file
$ cat stmt_file | mysql [database]</pre>
```

Creating a database

- Only database users with significant privileges can create databases
 - From the command line
 - \$ mysqladmin [opt] create dbname
 mysqladmin has the same command line options as mysql
 - From within the mysql monitor mysql> create database *dbname*

Exercises

- As root@localhost, create database 'biodb'
- Grant all privileges to the database user you created before
- Download the 1.sql (first take a look at the contents)
- Execute all SQL statements in the file

Hierarchy

A single MySQL service can have multiple databases
 mysql > SHOW databases;

• A particular database db can have multiple tables

```
mysql > USE db;
mysql > SHOW tables;
```

A particular table tbl can have multiple columns or fields

```
mysql > SHOW columns FROM tbl;
mysql > SHOW create table tbl;
```

Exercises

- Connect to the database service as a normal user
- What databases do you see?
- What tables are defined in biodb?
- What are the column names?

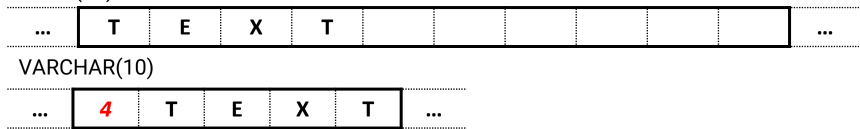
Relational databases with MySQL

- MySQL database = collection of tables
 - Table = set columns with specific types
 - Number, text, date, ...
 - Each row in same format
 - Only 1 value per field

Student_number	Name	Last_name	Birthdate	Sex
0293826	John	Doe	1991-10-02	M
0293749	Mel	Trotter	1991-04-11	V
0328273	Bill	Schuette	1990-12-01	M

- INT
 - Integer
 - SIGNED: -2 147 483 648 tot 2 147 483 647
 - UNSIGNED: 0 tot 4 294 967 295
 - TINYINT, BIGINT, SMALLINT
- FLOAT & DOUBLE
 - Numbers with decimal point
 - FLOAT: 7 digits after decimal point, DOUBLE: 15 digits after decimal point
- DATE
 - YYYY-MM-DD
 - DATETIME
 - YYYY-MM-DD HH:MM:SS
 - ! TIMESTAMP! No dates < 1970 and > 2038

- VARCHAR & CHAR
 - String with a certain number of characters
 - Define max number of characters e.g. VARCHAR(200)
 - VARCHAR: up to 65 535 characters
 - CHAR: up to 255 characters, spaces are added to reach required length CHAR(10)



- VARCHAR is more efficient in storage, CHAR is faster for reading data
- Similar for INT vs BIGINT vs ...

- TEXT & BLOB
 - Used for texts that are not queried often or do not have to be searchable
 - BLOB for binary data (images, ...)
- ENUM
 - List of permitted values
 - E.g. Set of colours: 'red', 'green', 'blue'
 - Very efficient

Student_number	Name	Last_name	Birthdate	Sex
0293826	John	Doe	1991-10-02	M
0293749	Mel	Trotter	1991-04-11	V
0328273	Bill	Schuette	1990-12-01	M

Column types

- Every row has to be unique
 - DBMS is able to distinguish separate rows

Student_number	Name	Last_name	Birthdate	Sex
0293826	John	Doe	1991-10-02	M
0293749	Mel	Trotter	1991-04-11	V
0328273	Bill	Schuette	1990-12-01	M

Primary key: column that makes sure every row is unique!

- Usually first column
- INT
- auto_increment: adds 1 to each value automatically

Constraints

On top of column types, there are some additional requirements per column

- Primary key
 - Only 1 PK per table, all values must be unique
- UNIQUE
 - All values (or combinations) must be unique
- NOT NULL
 - Field can not be empty when adding data (empty = null)
- Default
 - Default value for a field
- Foreign key
 - Same constraints as referenced column
 - Security when adjusting linked data possible

Summary – database structure

```
MySQL database

→ table

→ row

→ column

column type

constraint(s)
```

INSERT – add new rows

```
INSERT INTO tbl (col1, col2) VALUES (val1, val2);
INSERT INTO students (student_number, name, last_name) VALUES
(2654897, 'Glenn', 'Walker');
```

- Be aware!
 - Not all columns need to be included in query, unmentioned columns are given the default value for that column
 - Empty column gets null value
 - Columns with NOT NULL constraint require a value
 - Strings are written between quotes

SELECT – retrieve rows

```
SELECT columns FROM tbl;
SELECT * FROM modorg;
SELECT genus, species FROM modorg;
```

• columns

- List of columns, separated by comma
- * for all columns
- Use of arithmic operators (+, -, ...) and other functions (min, max, ...) on columns is possible

tbl

- Single table or multiple tables joined together
- Subquery
- View

ORDER BY-sort rows

```
SELECT columns FROM tbl ORDER BY col1 [asc|desc] [, col2 [asc|desc]...];
```

When using SELECT statements, the data is displayed in no particular order

→ use ORDER BY clause

- colX: a column or a column alias
- asc: ascending order (default)
- *desc*: descending order

Exercices

- Show the names (genus & species) of all model organisms in the order of the publishing date of the draft
- Show the names (genus & species) of all model organisms sorted by the number of chromosomes (most chromosomes on top) and than alphabetically by name

Calculated rows

- You can add columns in a query that calculate some value using other columns of the same row
- Lot of functions and operators readily available

```
mysql> SELECT 6*7;
mysql> SELECT concat(class, " ", genus) FROM modorg;
mysql> SELECT now();
```

Calculated rows – numbers

```
Operators+, -, *, /, %
```

Functions

```
sqrt(x), power(x, y), ...
exp(x), ln(x), ...
sin(x), cos(x)
round(x), ceil(x), floor(x), ...
rand(), rand(x)
```

Calculated rows – strings

```
• Functions
    length(s)
    concat(s1, ...)
    upper(s), lower(s)
    trim(s), ltrim(s), rtrim(s)
    substr(s, ...)
    reverse(s)
    truncate(s)
```

Calculated rows – dates

Functions

```
currentdate(), now()
year(d), month(d), week(d)
dayofmonth(d), dayofweek(d)
hour(d), minute(d), second(d)
```

Exercices

- Show
 - Model organism full name (as one column)
 - Genome size (as Gb), rounded to 4 digits
 - Average chromosome size
 - Publication year
 of all rows sorted by average chromosome size (largest on top)

Column aliases

- Columns can be renamed SELECT col [AS] alias ...
- The aliases can be used in the ORDER BY clause

Exercices

- Show
 - Model organism full name (as one column) as name
 - Average chromosome size as avgsize
 - Publication year as pubyear
 of all rows sorted by avgsize (largest on top)

WHERE- filter rows

SELECT columns FROM tbl WHERE condition(s) [ORDER BY sortcol];

- conditions
 - One or more conditions, combined with AND, OR, NOT, XOR
 - Only row for which the condition(s) evaluates TRUE are selected
 - Unable to use column aliases in condition(s)

Filtering rows – conditions

- Numerical comparison operators
 - =
 - != or <>
 - <, <=, >, >=
 - between x and y (inclusive)
- E.g. select all organisms with more than 10 chromosomes SELECT genus, species FROM modorg WHERE nchr > 10;

Filtering rows – conditions

- String comparison operators
 - =
 - != or <>
 - <, <=, >, >= (lexical)
 - like "pattern" matches a pattern
 - _ (A single character)
 - % (zero or more characters)
 - rlike "regex" [MySQL] matches a regular expression
- E.g. select all mammals

 SELECT genus, species FROM modorg WHERE class = "mammals";

Filtering rows – conditions

- Dealing with NULL-values
 - Testing for NULL-ness

```
SELECT ... WHERE col IS NULL;
SELECT ... WHERE col IS NOT NULL;
```

Substitution of NULL-values

```
SELECT ifnull(col, value) ... this function returns
```

- col if col is NOT NULL
- value if col is NULL

```
e.g. SELECT genus, species, ifnull(nchr, 0) FROM modorg;
```

Filtering rows – conditions

- Boolean logic
 - not x
 Evaluates TRUE if x is FALSE
 - x and y Evaluates TRUE if both x and y are TRUE
 - x or y Evaluates TRUE if x or y is TRUE, or both
 - $x \times x$ $y \in x$ $ext{x}$ $ext{clusive } ox$ $ext{or } y$ is TRUE, but not both

Exercises

- Select all mammals with genomes published after 2005
- Select all organisms that have an average chromosome size between 10 and 100 Mbp
- Select all organisms whose genus starts with A, B, C, D, or E

Filtering rows – duplicates

- Eliminate duplicate rows SELECT DISTINCT(cols) FROM ...
 - \rightarrow Each combination of cols is unique

Filtering rows – limiting output

- Limit the number of rows in a result set SELECT ... LIMIT n [OFFSET r];
- Result set is limited to a maximum of n rows
- ullet If an offset r is given, the first r rows are skipped
- Mostly used in combination with ORDER BY

Exercises

- Give an overview of all organism classes in the dataset (sorted alphabetically)
- Show the organism names of the top 3 largest genome sizes

Aggregation

- Queries are concentrated on particular rows
- Possible to calculate a single result across multiple rows e.g. maximum genome size?
- SQL allows you to
 - Specify criteria to group rows together
 - Calculate a single value per group
 - Filter grouped data

Aggregation

- Functions
 - count(*col*), count(*), count(distinct *col*)
 - sum(*col*)
 - min(col), max(col)
 - avg(col), stddev(col), variance(col)

Exercises

All these queries return a row count. What is the result and why?
 SELECT count(*) FROM modorg;
 SELECT count(nchr) FROM modorg;
 SELECT count(class) FROM modorg;
 SELECT count(DISTINCT class) FROM modorg;

How many mammals are in the database?

GROUP BY - aggregation

- Sort data into groups for aggregation purposes
 SELECT [col,] aggregatefunctions FROM src [WHERE cond] GROUP BY col [ORDER BY ...];
- All rows with the same value in col are grouped
- For each group, the aggregate function is calculated
- No sense in select other columns than col

Exercises

- How many organisms are present in the dataset for each class?
 Note the sort order.
- Show the minimum and the maximum genome sizes for each class.
 Take only those organisms into account for which genome sizes are known.
 Sort the results such that the biggest maximum genome size is on top.

Aggregation – filtering

• Filter results based on the results of aggregate functions using HAVING clause SELECT [col,] aggregatefunctions FROM src [WHERE cond1] GROUP BY col HAVING cond2 [ORDER BY ...];

• Column aliases can be used in *cond2*

```
SELECT
   FALL | DISTINCT | DISTINCTROW ]
     [HIGH PRIORITY]
     [STRAIGHT JOIN]
     [SQL_SMALL_RESULT] [SQL_BIG_RESULT] [SQL_BUFFER_RESULT]
      select_expr [, select_expr ...]
    [FROM table references
    [WHERE where condition]
    [GROUP BY { col_name | expr | position}]
     [ASC | DESC], ... [WITH ROLLUP]]
    [HAVING where condition]
    [ORDER BY { col_name | expr | position}]
     FASC | DESC], ...]
   [LIMIT {[offset,] row_count | row_count OFFSET offset}]
    [PROCEDURE procedure_name(argument_list)]
    [INTO OUTFILE 'file_name'
       [CHARACTER SET charset name]
       export options
       INTO DUMPFILE 'file_name'
       INTO var_name [, var_name]]
   [FOR UPDATE | LOCK IN SHARE MODE]]
```

Execution order

- 1. Input columns are determined
- 2. WHERE input columns are filtered
- 3. GROUP BY sorting & grouping of filtered input
- 4. Aggregation functions are calculated
- 5. HAVING aggregation functions are filtered
- 6. ORDER BY output is sorted
- 7. LIMIT/OFFSET output is chopped

Exercises

• For each class with more than 1 organism, show the average number of chromosomes. Sort the result such that the biggest average is on top.

Database upgrade

- Download 2.sql
- Create a new database bioinf
 mysql> CREATE database bioinf;
- Create the tables and insert the data
 \$ mysql bioinf < 2.sql

Joins

- Relation databases model entities and their relationships
- Different entities: different tables
- Allow you to combine information across different tables

Joins

· What if we want to expand our database with a trajectory and course info

Student_number	Name	Last_name	Birthdate	Trajectory	Course
0293826	John	Doe	1991-10-02	FBT	Databases
0293749	Mel	Trotter	1991-04-11	MLT	Databases
0328273	Bill	Schuette	1990-12-01	MLT	Databases
0293826	John	Doe	1991-10-02	FBT	Scripting
0293826	John	Doe	1991-10-02	FBT	Linux

- Problem: redundant information
 - Waste of space
 - Error prone

Joins

Solution: relational databases with a foreign key

Students

Student_number	Name	Last_name	Birthdate	Trajectory_ID	Course
0293826	John	Doe	1991-10-02	1	Databases
0293749	Mel	Trotter	1991-04-11	2	Databases
0328273	Bill	Schuette	1990-12-01	2	Databases
0293826	John	Doe	1991-10-02	1	Scripting
0293826	John	Doe	1991-10-02	1	Linux

 Important: Data types of linked columns have to be equal! Foreign key is typically primary key of other table

Trajectories

ID	Trajectory
1	FBT
2	MLT

Joins

Students

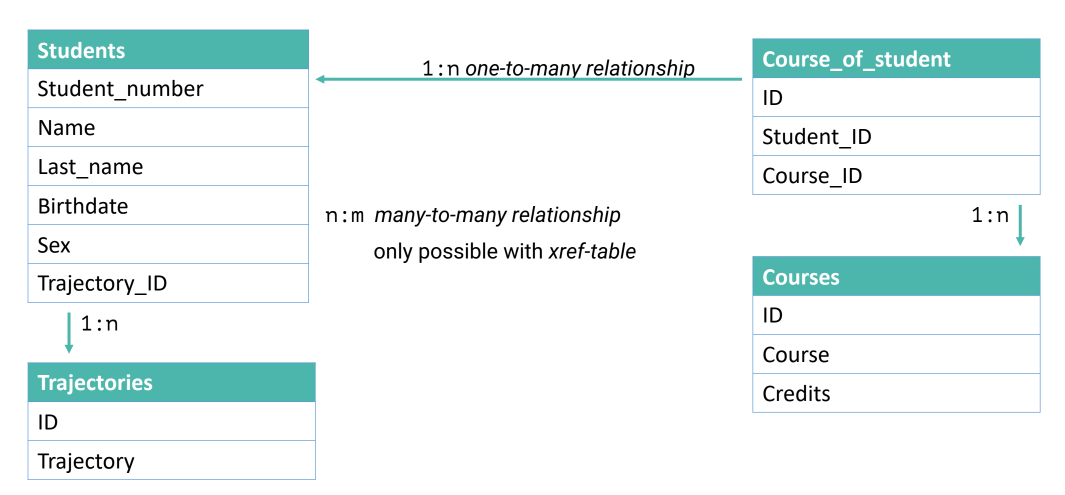
Student_number	Name	Last_name	Birthdate	Trajectory_ID
0293826	John	Doe	1991-10-02	1
0293749	Mel	Trotter	1991-04-11	2
0328273	Bill	Schuette	1990-12-01	2

Course_of_student

Student_id	Course_ID
0293826	1
0293826	2
0293826	3
0293749	1
0328273	1

ID	Course
1	Databases
2	Scripting
3	Linux

Joins



Database model= Entity Relationship Diagram (ERD)

Joins – Information is spread across multiple tables

- Create script that retrieves tables separately
- JOIN tables with query

Trajectory

Students		Student number	Name	Last_name	Birthdate	Course
Student_number		0293826	John	Doe	1991-10-02	Databases
Name		0293749	Mel	Trotter	1991-04-11	Databases
Last_name		0328273	Bill	Schuette	1990-12-01	Databases
Birthdate		0293826	John	Doe	1991-10-02	Scripting
Sex		0293826	John	Doe	1991-10-02	Linux
Trajectory_ID						
Trajectories	Data representation does not have to be equal to the the data is stored				the way	
ID						

Joins

Retrieve linked rows from different tables with JOIN

```
mysql> SELECT * FROM Students

JOIN Trajectories ON Students.Trajectory_ID = Trajectories.ID;
```

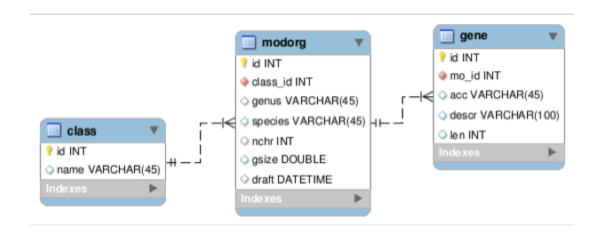
Students

Student_number	Name	Last_name	Birthdate	Trajectory_ID
0293826	John	Doe	1991-10-02	1
0293749	Mel	Trotter	1991-04-11	2
0328273	Bill	Schuette	1990-12-01	2
0324312	Penelope	Tracy	1989-07-24	NULL

ID	Trajectory
1	FBT
2	MLT
3	TI

Joins – back to our database

- modorg.class_id is a foreign key that references class.id
- gene.mo_id is a foreign key that references modorg.id



Joins – Cartesian product

Multiple tables in a query → database server generates all possible combinations
 = Cartesian product

table class has 6 rows
table modorg has 10 rows
query SELECT * FROM modorg, class has 60 rows

Joins

• Filtered Cartesian product
 SELECT * FROM modorg, class WHERE modorg.class_id = class.id;

SELECT * FROM modorg [INNER] JOIN class ON modorg.class_id = class.id;

Joins

Avoid ambiguity

```
mysql> SELECT id, name, genus, species FROM modorg, class WHERE
modorg.class_id = class.id;
```

```
ERROR 1052 (23000): Column 'id' in field list is ambiguous
```

Ambiguous columns must be qualified. And additionally you can choose an alias:
 mysql> SELECT modorg.id as mo_id, name, genus, species FROM modorg, class WHERE modorg.class_id = class.id;

Joins – data source alias

- Join table with itself or select data from subquery
 - Use alias for data source

```
mysql> SELECT a.col, b.col FROM src1 [as] a, src2 [as] b WHERE ... mysql> SELECT a.col, b.col FROM src1 [as] a JOIN src2 [as] b ON ...
```

Joins – data source alias

 For each class, give the class name, organism name and date of the organism that was sequenced first

```
mysql> SELECT class_id, min(draft) as dr FROM modorg GROUP BY
class_id;
```

- add class name: join with table class
- add organism name: join with table modorg

Joins - data source alias

Add class name

```
mysql> SELECT name, dr FROM
    (SELECT class_id, min(draft) as dr FROM modorg GROUP BY
    class_id) as s, class WHERE s.class_id=class_id;
```

add organism name: join with table modorg

```
mysql> SELECT name, concat(genus," ",species) as org_name, dr
   FROM (SELECT class_id, min(draft) as dr FROM modorg GROUP BY
   class_id) as s, class, modorg WHERE s.class_id=class.id AND
   s.dr=draft;
```

Joins – 4 types

- INNER JOIN
 - Only rows present in both tables
- LEFT JOIN
 - All rows from left table, even without linked data in right table
- RIGHT JOIN
 - All rows from right table, even without linked data in left table
- OUTER JOIN
 - All rows from both tables
 - Doesn't exist in MySQL

Joins - INNER JOIN

Students

Student_number	Name	Last_name	Birthdate	Trajectory_ID
0293826	John	Doe	1991-10-02	1
0293749	Mel	Trotter	1991-04-11	2
0328273	Bill	Schuette	1990-12-01	2
0324312	Penelope	Tracy	1989-07-24	NULL

ID	Trajectory
1	FBT
2	MLT
3	Ti



Student_number	Name	Last_name	Birthdate	Trajectory
0293826	John	Doe	1991-10-02	FBT
0293749	Mel	Trotter	1991-04-11	MLT
0328273	Bill	Schuette	1990-12-01	MLT

Joins - LEFT JOIN

Students

Student_number	Name	Last_name	Birthdate	Trajectory_ID
0293826	John	Doe	1991-10-02	1
0293749	Mel	Trotter	1991-04-11	2
0328273	Bill	Schuette	1990-12-01	2
0324312	Penelope	Tracy	1989-07-24	NULL

ID	Trajectory
1	FBT
2	MLT
3	Ti



Student_number	Name	Last_name	Birthdate	Trajectory
0293826	John	Doe	1991-10-02	FBT
0293749	Mel	Trotter	1991-04-11	MLT
0328273	Bill	Schuette	1990-12-01	MLT
0324312	Penelope	Tracy	1989-07-24	NULL

Joins - RIGHT JOIN

Students

Student_number	Name	Last_name	Birthdate	Trajectory_ID
0293826	John	Doe	1991-10-02	1
0293749	Mel	Trotter	1991-04-11	2
0328273	Bill	Schuette	1990-12-01	2
0324312	Penelope	Tracy	1989-07-24	NULL

ID	Trajectory
1	FBT
2	MLT
3	TI



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NULL	NULL	NULL	NULL	TI

Joins - OUTER JOIN

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Joins – characteristics

- * is used to select all columns from all tables
- Use *tbl.col* to specify column
- Very inefficient
 - A lot of memory, a lot of time
- Use ON to specify linked columns

Execution order

- 1. Input columns are determined
 - a. JOIN clause
- 2. WHERE input columns are filtered
- 3. GROUP BY sorting & grouping of filtered input
- 4. Aggregation functions are calculated
- 5. HAVING aggregation functions are filtered
- 6. ORDER BY output is sorted
- 7. LIMIT/OFFSET output is chopped

Exercises

- Experiment with the different types of JOINs
- For all rows in table gene, show
 - Organism name
 - Class name
 - Accession
 - Length
 - Description of the gene

Efficiency and speed

- Complex queries tend to become slow
 - Large system load
 - Slow interface, user has to wait
- Mind your choices of column types and included columns in the query
- Frequently used queries can be sped up
 - Views
 - Indices
 - Allow redundancy

Views

- Re-use same query
- Query can be saved as a special table-like object
- Usually read-only data source
- Speed gain depends on use case
- MySQL
 - Virtual table
 - Used to serve up data in an orderly fashion
- Oracle i.a.
 - Materialized view: result of a SELECT query is stored
 - Very efficient

Views

Create a new view
 mysql> CREATE VIEW viewname as SELECT ...

Use a view as a table
 mysql> SELECT ... FROM viewname WHERE ... ORDER BY ...

Exercises

- Create a view (genevw) from the query in the previous exercise
- Select all genes containing hemoglobin in the description and sort the result set by gene length.

Next:

- What is the minimum and maximum gene length?
- What is the average gene length?
- And the standard deviation?
- Does the view show up when using

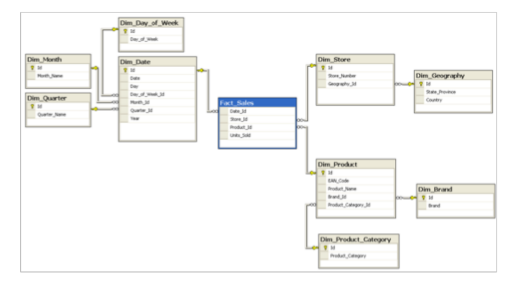
```
mysql> show tables;
```

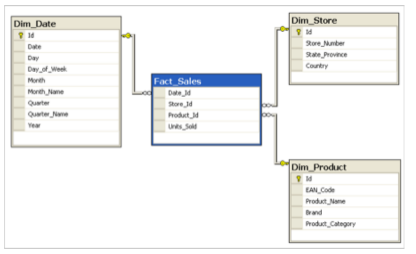
Index

- Quickly find certain rows
- Stored separately
- Golden rule: Use indices on columns you will use in your WHERE clause
- Only 1 index per query is used
 CREATE [UNIQUE|FULLTEXT|SPATIAL] INDEX index_name ON tbl (col);
- Foreign key
 - Index used to speed up JOIN queries
 - Not essential for JOIN queries

Allow redundancy

- Schema with least redundancy isn't always the quickest
- Allow redundancy to reduce number of JOINs
- E.g. Data warehouse with real-time reporting
 - High efficiency is required
- Snowflake vs Star schema





Snowflake

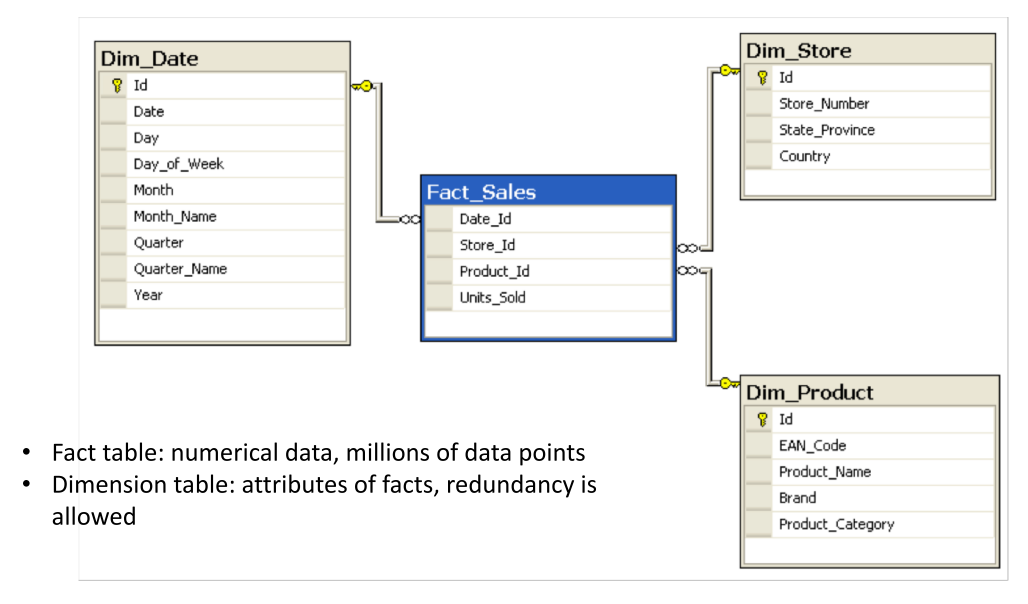
Star

P.Product Category Id = C.Id

GROUP BY B.Brand, **G**.Country

WHERE D.Year = 1997 AND C.Product Category = 'tv'

SNOWFLAKE STAR Redundant data No redundancy Easy to maintain and change Less easy to maintain/change Complex queries Lower query complexity Slower (more JOINs) **Faster** Uses less space Uses more space (data is stored twice or more) Bottom up Top down SELECT B.Brand, SELECT P.Brand, G. Country, S.Country AS Countries, **SUM**(F.Units Sold) **SUM**(F.Units Sold) FROM Fact Sales F FROM Fact Sales F **INNER JOIN** Dim Date D **ON** F.Date Id = D.Id **INNER JOIN** Dim Date D **ON** (F.Date Id = D.Id) **INNER JOIN** Dim Store S **ON** F.Store Id = S.Id **INNER JOIN** Dim Store S **ON** (F.Store Id = S.Id) **INNER JOIN** Dim Geography **G ON** S.Geography Id = **G**.Id INNER JOIN Dim_Product P ON (F.Product_Id = P.Id) INNER JOIN Dim Product P ON F. Product Id = P.Id WHERE D.Year = 1997 AND P.Product_Category = 'tv' **INNER JOIN** Dim Brand B **ON** P.Brand Id = B.Id **GROUP BY** P.Brand, S.Country INNER JOIN Dim Product Category C ON



Database backup

- Dump a complete database into a tekst file
 - \$ mysqldump [opt] db > db.sql
- Includes
 - Statements for creating the database if the option --databases is used
 - Statements for creating tables, views, ...
 - Statements for inserting data
- Restore the database (may need to create first)
 - \$ mysql db < db.sql</pre>

Exercises

- Create a separate dump file for each of your own databases
- Check the contents of each file
- For the bold ones
 - Drop your databases and recreate them using your dump files