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1. Introduction

This is the manual for the Leiden Open (source) Variation Database (LOVD) version 3.0. LOVD 3.0 is a partial rewrite of LOVD 2.0, which first stable release was completed in 2007. Also, LOVD 3.0 has a greatly improved database model, and includes lots of new features aimed at making LOVD useful for more research environments.

LOVD is designed to provide a flexible, freely available tool for gene-centered collection and display of DNA variants. LOVD 3.0 extends this idea to also provide patient-centered data storage and storage of NGS data, even of variants outside of genes.

LOVD was developed approaching the “LSDB-in-a-Box” idea for the easy creation and maintenance of a fully web-based gene sequence variant database, that is platform-independent and uses PHP and MySQL open source software only. The design of the database follows the recommendations of the Human Genome Variation Society (HGVS) and focuses on the collection and display of DNA sequence variants, but it has fully implemented methods for storing complete clinical data as well. The open LOVD setup also facilitates functional extensions with scripts written by the community.

The development of (then nameless) LOVD started in late 2002, while it was first officially released in January, 2004. Before that LOVD was only in use by the Leiden Muscular Dystrophy pages, as a not-so-modular system with lots of characteristics specific for that website only. With the official release of LOVD in 2004 the system had become much more dynamic and customizing LOVD was made easy mostly by editing text-files.

In 2004, LOVD became available under the open source license GPL and with the 1.1.0 release most of the text-files had been replaced by online forms so customizations can be performed through the web interface. Early in 2005 the first LOVD article was published, and in 2005 the development of LOVD was more targeted at improving the ease of use of the system.

In 2006 the development of LOVD 2.0 started after the decision was made to rewrite all of LOVD from scratch to be able to include a long list of upgrade suggestions that were hard to implement in LOVD 1.1.0. Aimed at modularity and data redundancy, LOVD 2.0 was meant to be a more flexible and more powerful successor of the popular 1.1.0 version and soon it received the interest of LOVD users eager to try out the all-new version.

With more features being added and bugs fixed rapidly, LOVD 2.0 reached beta stage in April 2007, after which more and more users started to upgrade their 1.1.0 databases to 2.0. Finally, in October 2007 LOVD 2.0 reached the stable stage, after which LOVD 2.0 was continuously improved with monthly releases for two years, after which the releases became less frequent. LOVD 2.0 is described in the second LOVD paper.

By 2009 it had became clear that although LOVD 2.0 was a great step forward, there were still key improvements to be made. Since the complexity of necessary changes had become to great to gradually upgrade LOVD 2.0 systems to include these options, it was again decided to start from scratch writing LOVD 3.0. This allowed us to redesign the complete data model in full freedom, although it should still be possible for existing LOVD 2.0 databases to have all data transferred to LOVD 3.0.

LOVD 3.0 adds even more flexibility, allowing users to focus exclusively on sequence variants, whilst also allowing an exclusive focus on individuals and clinical data, and anything in
1. Introduction

between. It will be possible for different submitters to work together cooperatively on the same data. Searching through the data is improved extensively, and webservises of many different sources are used to automatically retrieve gene and transcript information. Also new in 3.0 is full Next Generation Sequencing (NGS) support, with the ability to import VCF files. For VCF file imports, LOVD allows for automatic annotation of the variants. Both formats support the automatic creation of genes and transcripts in the system, greatly reducing the amount of work required by curators to get LOVD set up for their research data.

LOVD 3.0 reached beta stage in January 2012, and the first stable release was in December of that year. Currently, the latest release is 3.0-30. Keep an eye on our news page for the latest information on LOVD 3.0 development.

Wherever you see “he” or “his” written in this manual, it should read “he or she” and “his or her” respectively.

Please note that this manual is work in progress. Since LOVD 3.0 is still under development and the development is the focus of our efforts, many features in LOVD 3.0 are not yet described in this manual. Also, features described in this manual may become inaccurate or even incorrect in later versions of LOVD 3.0. Please bear with us while we finish this manual.
2. View and search data in an LOVD installation

To navigate in an LOVD installation you can use the tabs on the top of the page. Which tabs you can see depends on whether you are logged in or not, and your user level. When you are not logged in or you are logged in as a submitter, you see the following tabs: Genes, Transcripts, Variants, Individuals, Diseases, Screenings, Submit and Documentation. Additionally, Curator-level users and up can see the Configuration tab and only Managers and the Database administrator can see the Users and Setup tabs.

Each tab has a dropdown menu and the tab itself has a default page to which you will go when you click on the tab. The options of the dropdown menus depend on your user level.

2.1 Searching and sorting

All list views have search fields which can be used to search data. You can search for a complete word or you can search for a part of a search term. For example, when you search for “AAM” in the Symbol field of the genes listing, LOVD will not only find genes starting with “AAM”, but also containing “AAM”, like “DAAM1” and “DAAM2”. To search for prefixes or suffixes, see the table further below.

To sort on a certain column, click on the column header or on the arrows. If that column is already selected to sort on, the sort order will be swapped. The column currently sorted on has a darker blue background color than the other columns. The up and down arrows next to the column name indicate the current sorting direction. When sorting on any field other than the default, LOVD will sort secondarily on the default sort column.

LOVD allows boolean search terms, meaning you can construct complex queries with AND, OR and NOT logic. Typing a space between search terms is treated as AND, OR is indicated with a pipe ‘|’ with no spaces around it and NOT is indicated by an exclamation mark ‘!’, not followed by a space. If you enclose two or more words in double quotes, LOVD will search for the combination of those words only exactly in the order you specify. Note that search terms are case-insensitive and that wildcards such as * are treated as normal text!
You can use different operators to specify your search request:

<table>
<thead>
<tr>
<th>Operator</th>
<th>Example</th>
<th>Matches</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arg</td>
<td>Results containing ‘Arg’.</td>
<td></td>
</tr>
<tr>
<td>Arg Ser</td>
<td>Results containing both ‘Arg’ and ‘Ser’.</td>
<td></td>
</tr>
<tr>
<td>Arg</td>
<td>Ser</td>
<td>Results containing ‘Arg’ or ‘Ser’.</td>
</tr>
<tr>
<td>!fs</td>
<td>Results not containing ‘fs’.</td>
<td></td>
</tr>
<tr>
<td>p.(Arg)</td>
<td>Results beginning with ‘p.(Arg’ (prefix search).</td>
<td></td>
</tr>
<tr>
<td>Ser)$</td>
<td>Results ending with ‘Ser)’ (suffix search).</td>
<td></td>
</tr>
<tr>
<td>!&quot;&quot;</td>
<td>Results with this field empty.</td>
<td></td>
</tr>
<tr>
<td>!='p.0'</td>
<td>Results exactly matching ‘p.0’ (exact search).</td>
<td></td>
</tr>
<tr>
<td>&quot;p.0&quot;!</td>
<td>Results with this field not empty.</td>
<td></td>
</tr>
<tr>
<td>!&quot;&quot;!</td>
<td>Results not exactly matching ‘p.0’.</td>
<td></td>
</tr>
<tr>
<td>*</td>
<td>Ter !fs</td>
<td>Results containing ‘*‘ or ‘Ter’ but not containing ‘fs’.</td>
</tr>
<tr>
<td>&lt;2020</td>
<td>Values before 2020 (date fields).</td>
<td></td>
</tr>
<tr>
<td>&lt;=2020-06</td>
<td>Values on or before June, 2020 (date fields).</td>
<td></td>
</tr>
<tr>
<td>&gt;2020-06</td>
<td>Values after June, 2020 (date fields).</td>
<td></td>
</tr>
<tr>
<td>&gt;=2020-06-15</td>
<td>Values on or after June 15th, 2020 (date fields).</td>
<td></td>
</tr>
<tr>
<td>&gt;2009 &lt;2020</td>
<td>Values from 2010 to 2019 (date fields).</td>
<td></td>
</tr>
<tr>
<td>&lt;23</td>
<td>Values lower than 23 (numeric fields).</td>
<td></td>
</tr>
<tr>
<td>&lt;=23</td>
<td>Values lower than, or equal to, 23 (numeric fields).</td>
<td></td>
</tr>
<tr>
<td>&gt;23</td>
<td>Values higher than 23 (numeric fields).</td>
<td></td>
</tr>
<tr>
<td>&gt;=23</td>
<td>Values higher than, or equal to, 23 (numeric fields).</td>
<td></td>
</tr>
<tr>
<td>&gt;=20 &lt;30 !23</td>
<td>Values from 20 to 29, but not equal to 23 (numeric fields).</td>
<td></td>
</tr>
</tbody>
</table>

Some more advanced examples:

- **Asian**  Results containing ‘Asian’, but also containing ‘Caucasian’.
- **Asian !Caucasian**  Results containing ‘Asian’, but not containing ‘Caucasian’.
- **"South Asian"**  Results containing ‘South Asian’, but not ‘South East Asian’.

## 2.2 Genes

When no gene is selected, the default page of the genes tab shows all genes. When a gene is selected (shown in the page header), the default page of the gene tab is the gene homepage.

**View all genes**

Here you can see all the genes for this installation. When you click on a gene entry, you’re taken to the gene homepage.

**View gene homepage**

Here you can see the details of a gene. For more information, see section “The gene homepage”. This menu item is not available when no gene has been selected.

**View graphs**

This displays the statistics of the variants within a gene. This menu item is not available when no gene has been selected.
Create a new gene entry
This menu item is only available for Managers or higher. See section “Creating a new gene database” on how to add a new gene.

2.3 Transcripts

When no gene is selected, the default page of the transcripts tab shows all transcripts. When a gene is selected (shown in the page header), the default page of the transcripts tab shows all transcripts of the selected gene.

View all transcripts
View all transcripts stored in the database. Click on the transcript to view the transcript’s details and variant overview.

View all transcripts of a gene
View all transcripts of the currently selected gene. Click on the transcript to view the transcript’s details and variant overview. This menu item is not available when no gene has been selected.

Create a new transcript information entry
This menu item is only available for Curators or higher. See section “Creating new transcripts” on how to add a new transcript to a gene.

Transcript detailed view
When clicking a transcript from one of the transcript data listings, the transcript’s details are shown. Below the transcript’s details, a variant listing is shown of all variants found on this transcript. Click on a variant to view the variant’s details.

2.4 Variants

When no gene is selected, the default page of the variants tab shows all variants on the genome level. When a gene is selected (shown in the page header), the default page of the variants tab shows all unique variants in the selected gene.

View all variants
This view displays all variants in this LOVD installation on the genome level. Variants within, but also outside of genes are shown. You can click on a variant to view its details.

View all variants affecting transcripts
This view displays all variants in this LOVD installation that are found within a gene, and have been mapped to a transcript. You can click on the a variant to view its details.
2. View and search data in an LOVD installation

**View unique variants in a gene**
This view displays the gene’s unique variants. Each variant is shown only once, regardless of the number of observations. The number of times a variant is found is displayed in the field “Reported”. The information in other fields are summarized and not all information for the variants may be displayed. Click on a variant to see all reported observations. This menu item is not available when no gene has been selected.

**View all variants in a gene**
View of all variants reported in the selected gene. Variants are not grouped like in the unique variant view. Click on a variant to see the variant’s details. This menu item is not available when no gene has been selected.

**Full data view for a gene**
This view shows all variant observations together with the variant screening(s) and the individual in which the variant has been reported. Although this data view is very wide, it allows searching for variants only found in individuals with certain characteristics. Click on a variant to see the variant’s details. This menu item is not available when no gene has been selected.

**Create a new data submission**
This menu item is only available when logged in. See chapter “Adding data to LOVD” on how to create a new data submission.

**View active genomic custom columns**
This menu item is only available for Managers or higher. See chapter “Custom columns” for more information on custom columns.

**Enable more genomic custom columns**
This menu item is only available for Managers or higher. See section “Enabling custom columns” on how to enable custom columns.

**Variant detailed view**
When clicking on a variant from one of the variant data listings, its details are shown. If the variant is affecting any transcripts, a listing is shown below the variant’s genomic details. Click on any of the entries to view the data in a tabular view. If the variant has been found in an individual, at the bottom of the page the listing for the variant screenings used to identify the variant is shown. On the top of the page a link is shown to the individual in which the variant is found.

### 2.5 Individuals

When no gene is selected, the default page of the individuals tab shows all individuals. When a gene is selected (shown in the page header), the default page of the individuals tab shows all individuals with variants in the selected gene.

**View all individuals**
This view shows all individuals in this LOVD installation. You can click on an individual to view its details.

![Figure 2.5: Individuals menu.](image)
2. View and search data in an LOVD installation

View all individuals with variants in a gene
This view displays all individuals with variants in the currently selected gene. You can click on an individual to view its details. This menu item is not available when no gene has been selected.

Create a new data submission
This menu item is only available when logged in. See chapter “Adding data to LOVD” on how to create a new data submission.

View active custom columns
This menu item is only available for Managers or higher. See chapter “Custom columns” for more information on custom columns.

Enable more custom columns
This menu item is only available for Managers or higher. See section “Enabling custom columns” on how to enable custom columns.

Individual detailed view
When clicking on an individual from one of the data listings, the individual’s details are shown. Below the individual’s details, the phenotype entries are shown if any are stored. Also variant screenings performed are listed, and variants found in this individual. Clicking on any of these entries will show the respective details.

2.6 Diseases
When no gene is selected, the default page of the diseases tab shows all diseases. When a gene is selected (shown in the page header), the default page of the diseases tab shows all diseases associated with the selected gene.

View all diseases
This view shows all the diseases in the database. Also the number of individuals reported with this disease is displayed and the number of phenotypes reported. You can click on a disease to view its details.

View all diseases associated with a gene
This view displays all diseases associated with the currently selected gene. Also the number of individuals reported with this disease is displayed and the number of phenotypes reported. You can click on a disease to view its details. This menu item is not available when no gene has been selected.

Create a new data submission
This menu item is only available when logged in. See chapter “Adding data to LOVD” on how to create a new data submission.

View available phenotype columns
This menu item is only available for Curators or higher. See chapter “Custom columns” for more information on custom columns.
Disease detailed view
On the disease’s detailed view, all individuals diagnosed with this disease are listed. From the table, a link is provided to view all phenotype entries for this disease.

2.7 Screenings

When no gene is selected, the default page of the screenings tab shows all screenings. When a gene is selected (shown in the page header), the default page of the screenings tab shows all screenings that screened for variants in the selected gene.

View all screenings
This view shows all the screenings in the database. You can click on a screening to view its details.

View all screenings for a gene
This view displays all screenings that screened for variants in the currently selected gene. You can click on a screening to view its details. This menu item is not available when no gene has been selected.

Create a new data submission
This menu item is only available when logged in. See chapter “Adding data to LOVD” on how to create a new data submission.

View active custom columns
This menu item is only available for Managers or higher. See chapter “Custom columns” for more information on custom columns.

Enable more custom columns
This menu item is only available for Managers or higher. See section “Enabling custom columns” on how to enable custom columns.

Screening detailed view
On the screening’s detailed view, all genes screened for variants are listed, as well as any variants found.
3. Installing LOVD

Please note that if you’re just going to *use an LOVD installation that already exists*, you do not need to install anything on your computer. Any web browser that you already have installed, such as Mozilla Firefox, Google Chrome or Microsoft Internet Explorer, will do. This chapter describes how to install a new LOVD.

Installing LOVD is a “piece of cake”. However, you might require help from your system administrator, since there are a few dependencies that need to be taken care of first.

3.1 Before you install

LOVD is a web-based software project. Therefore, installation requires a correctly set-up webserver. A suitable server with the necessary software installed is available at virtually all academic institutions and countless commercial hosting providers. If you don’t want to run LOVD on a server but rather want to have it running on your computer, we can recommend the XAMPP package, which installs all needed software on your computer regardless of whether it’s Linux, Windows or Mac. For Windows users, a working alternative is the WampServer package. Also, at the time of this writing, we are offering free server space if you need to set up an LOVD. See our website for more information. In that case you don’t need to install anything and you can skip this chapter.

If you’re experienced with web servers and you want to set it up yourself: LOVD has been extensively tested with the Apache webserver, but any webserver able to run PHP scripts should suffice. You’ll need PHP 5.3.0 or higher with mbstring support enabled and openssl installed and MySQL 4.1.2 or higher; MariaDB also works as an alternative to MySQL. LOVD will force the use of the UTF-8 character set, but we recommend setting this as the default value for your software nonetheless. Also, you’ll need to install a properly configured Mail Transfer Agent for sending emails (such as registration information).

LOVD may work with other database platforms, but we are not developing for other database backends. You are welcome to try other database backends and report the results to us, but we can’t provide any support for it. For more information, see the FAQ about software needed.

Before installing LOVD, be sure you have the required credentials for connecting to the MySQL database. You need a hostname, username, password and a database name to be able to install LOVD. If you are installing LOVD on a remote server, be sure to have the (s)FTP username and password to be able to copy the necessary files to the server.

Download & Extract

To download LOVD, go to the LOVD website and click on the “Download” tab. You can download LOVD in two formats; GZIPped TARball and ZIP archive. The first format is common for Unix and Linux systems while the ZIP archive is popular on the Windows platform. Usually you will be able to open both formats.

Download the file of your choice and save it to your hard drive. Now extract the file using GZIP/TAR or ZIP to the desired location. On a server, common directories may
be /var/www on Unix or Linux servers, or C:\htdocs on a Windows server. When using XAMPP, the directory is C:\xampp\htdocs for Windows, /opt/lampp/htdocs for Linux, or /Applications/XAMPP/htdocs for Mac. To be sure, consult the documentation of the web-server software you are using.

Pre-install Setup

You will need to rename the standard config file config.ini.php-lovd to config.ini.php and edit it in, for example, a basic text editor. This is absolutely mandatory, because you will need to enter the MySQL hostname, username, password and database name here. In case you’re using XAMPP, the needed values are “localhost”, “root”, “” (empty) and “test”, respectively. No quotes around the values needed. Please note that these are not the recommended settings for production environments, but it will help you quickly set up LOVD on your computer.

Please go through the entire config.ini.php file to determine if you need to change any of the other settings.

A .htaccess file is put in the root directory of your LOVD installation protecting the config.ini.php file, as well as setting some PHP options. This will prevent the config file from being accessed by others on Apache HTTP servers (if configured properly), the most commonly used webserver. If you use Apache, please check that your version and configuration support this feature. Make sure you have the .htaccess file into your LOVD directory, on Unix and Linux systems it’s a hidden file so it can be missed easily. For the .htaccess file to work, you need to have “Limit” and “Options” enabled in Apache’s “AllowOverride” setting.

Also, make sure you have MultiViews or mod_rewrite enabled. This allows a PHP file like “/setup.php” to be accessed as “/setup”. The mod_rewrite rules in the .htaccess file require “FileInfo” enabled in Apache’s “AllowOverride” setting.

More information about .htaccess files:
http://httpd.apache.org/docs/2.0/howto/htaccess.html

More information about AllowOverride:
http://httpd.apache.org/docs/2.0/mod/core.html#allowoverride

If you use a different webserver, make sure to configure it to deny access to the config.ini.php file. LOVD will access the file through the filesystem. Also, since the .htaccess file sets a couple of options, please disable the following PHP options: register_globals, magic_quotes_gpc, mysql.trace_mode. Also make sure there is “MultiViews” functionality, which allows a PHP file like “/setup.php” to be accessed as “/setup”.

3.2 Install process

To install LOVD on a remote webserver, upload the LOVD directory with all the files to the webserver by, for instance, (s)FTP. If you install LOVD on your own computer, you do not need that step.

Next, point your web browser to the directory where you’ve saved the LOVD files. This could, for instance, be http://localhost/LOVDv.3.0/ or, if you’re using a remote server,
something like http://www.your-domain.org/LOVD/. LOVD will tell you it’s not installed yet, and include a link to the install directory.

Figure 3.1: When pointing your browser to the LOVD location, it will tell you it’s not installed yet. Click the link to start the install process.

LOVD will first check a few requirements. Both the PHP and MySQL versions will be checked, to make sure your LOVD will function properly on your webserver environment. Also some settings of the web server, PHP and MySQL will be checked. Finally, LOVD will check if your config file has been hidden. If all looks well, LOVD will tell you all requirements are OK and you’re ready to start installing LOVD.

Installing LOVD consists of only 4 simple steps and will take only a couple of minutes, or less.

Figure 3.2: If all requirements are OK, you’re ready to start installing LOVD.

Administrator account details

Fill in the database administrator data to install LOVD. The database administrator will be the first user registered with LOVD and has full access to all of LOVD’s functionalities. The database administrator is the only user capable of creating manager accounts. Managers can only create curator and submitter accounts.

Most of the form is pretty straightforward, but I will highlight one field - “Allowed IP address list”. An IP address is an address a computer is known by on the network. To help prevent others to try and guess the username/password combination, you can restrict access to the database administrator account to a number of IP addresses or ranges. This also means you need to be very careful with this setting, as being too restrictive may lock you out of your account. The default, unrestricted, value is *.
3. Installing LOVD

Installation process

After completing the database administrator account details, click the “Continue »” button at the bottom. LOVD will apply a simple username and password quality check. If LOVD tells you that the provided details are OK, click the “Next »” button.

Installing database tables

The next step is to create and fill all necessary LOVD database tables. LOVD will do this automatically, and you can watch the progress bar complete while the tables are created, the country list filled, the database administrator account created, the custom columns preconfigured, the standard columns enabled, and the custom links generated. This may take a while on non-optimal conditions, so please be patient. When everything is done, a “Next »” button appears - click it to continue.

Configuring LOVD system settings

The final form in the installation process is completing the initial configuration of the LOVD system settings. These settings can be changed after installation at any time through the
LOVD setup.

The only two settings that cannot be changed at a later time, are the install lock, which is checked by default, and the human build that the system will use to map your variants to. Setting the uninstall lock will prevent uninstallation of LOVD by the database administrator. The only way to remove this lock after installation is directly through the MySQL database backend.

For more information on the options of the LOVD system settings, see the LOVD system settings section of the manual.

After filling in the form, click “Continue ». If everything was filled in correctly, LOVD will register the LOVD system settings. Click “Next »” to continue to the last step. Almost done!
3. Installing LOVD

Install process

When everything has been filled in and stored correctly, the installation is complete. Now that you’re done installing LOVD, click the “Continue to Setup area” button to be forwarded to the LOVD setup area. From there, you can perform the most important actions for managers in LOVD, such as creating new gene databases or disease information entries, registering new user accounts or authorizing curators, or manage the custom columns and links. The button to create a new gene will be highlighted, as a suggestion of your next step!
4. LOVD Setup

To change system-wide settings, but also to perform actions with system-wide effects, start at the LOVD setup area. The setup area is only available for managers and the database administrator and can be accessed by clicking on the “Setup” tab in the menu. If you do not see a “Setup” tab, you first need to log into the system with a manager or administrator account.

The LOVD setup shows some statistics on the left of the screen, like date of installation, number of registered users and variant counts, separated by status. Furthermore, there are two columns of options. The left column has options for the system settings, authorized users, the custom columns, custom links, full data download and the LOVD system logs. The right column has options for genes, transcripts, diseases, individuals and variants. Some of the general options from the LOVD setup are also available through the Setup tab’s dropdown menu, allowing you to quickly navigate to other setup options without having to go through the setup main page. The other options shown in the setup area are available through their own tab’s dropdown menus.

In this chapter, we will describe changing the LOVD system settings, how to check, search and clean up the LOVD system logs, and how to uninstall LOVD. The other actions available from the LOVD setup are described in their respective chapters (see the table of contents).

4.1 LOVD system settings

The LOVD system settings allow you to adapt your LOVD installation to your preferences. For instance by changing the installation’s displayed name or email address (general settings), proxy server settings (connection settings), settings on statistics, or the security settings.

To view or edit the LOVD system settings, click on the “Setup” tab, then on “View and change LOVD System settings (...),” or move your mouse over the “Setup” tab and click the “LOVD system settings” menu option from the dropdown list.

General system settings

**Title of this LOVD installation**

This title will be shown on the top of every page, above the menu tabs. The default value is “LOVD - Leiden Open Variation Database”.

**Institute**

The institute which runs this database is displayed in the public area and in emails sent by LOVD. It’s commonly set to a laboratory name or a website name. If it’s not specified, LOVD will use the (autodetected) website name LOVD has been installed on.
Database URL
This is the URL with which the database can be accessed by the outside world, including “http://” or “https://”, and is not necessarily the URL you are using right now to access the database. It will also be used in emails sent by LOVD. This field is mandatory if you select the “Include in the global LOVD listing” option.
If you click the “check” link, LOVD will verify or try to predict the value.

LOVD email address
This email address will be used to send emails from LOVD to users. LOVD needs this address to prevent problems with spam filters, to make sure that emails from LOVD arrive. Please note that although strictly speaking this email address does not need to exist, we recommend that you use a valid address to make sure that bounces of emails sent from LOVD (to submitters or curators) are still caught by someone, so that they can be handled.

Forward messages to database admin
With this setting enabled, LOVD will forward messages to the database administrator about submitter registrations, submissions, and such.

Human Build to map to (UCSC/NCBI)
This value can only be set during installation, and cannot be changed later. It defines on which version of the Human Build (hg18/Build 36.1 or hg19/GRCh37) genomic variants are described. Links to the UCSC and Ensembl genome browsers and the NCBI sequence viewer also depend on this setting.
From LOVD 3.0-beta-07, the hg18 setting is no longer available for new LOVD installations, although it will remain to be supported. When hg20 will come out, LOVD will start supporting multiple genome builds per installation.

List database changes in feed for how long?
*Note: Not yet implemented!
LOVD includes a “newsfeed” that allows users to get a list of changes recently made in the database. Select here how many months back you want changes to appear on this list. Set to “Not available” to disable the newsfeed.

Connection settings (optional)
Some networks have no access to the outside world except through a so-called proxy server (more info on WikiPedia). If this applies to the network this server is installed on, please fill in the proxy server information here.

Proxy server host name
The host name of the proxy server, such as “www-cache.institution.edu”.

Proxy server port number
The port number of the proxy server, such as 3128.

Proxy server username
In case the proxy server requires authentication, please enter the required username here.

Proxy server password
In case the proxy server requires authentication, please enter the required password here.
Customize LOVD

Here you can customize the way LOVD looks. Right now, there is only one option included, we will add more later.

System logo
If you wish to have your custom logo on the top left of every page instead of the default LOVD logo, enter the path to the image here, relative to the LOVD installation path. If left empty, it will be set to the default value, which is “gfx/LOVD_logo130x50.jpg”. Currently, only images already uploaded to the LOVD server are allowed here.

Global LOVD statistics

Your LOVD installation can send some general information on your installation back to us. We use this information to see how many LOVDs there are worldwide, which versions of LOVD are currently in use, how LOVD is being used (amount of data), to see what type of software is used to run LOVD and to construct the public list of LOVD installs.

Send statistics
When this setting is enabled, LOVD will collect general usage statistics and send this to us. The following information is sent: the number of submitters, genes, individuals, variants and unique variants in your system. No specific information is sent, just the numbers.

Include in the global LOVD listing
On our website, we keep a list of public LOVD installations. This list is also feeding our worldwide LSDB list, through which submitters can locate databases for a specific gene to submit data to. If you enable the “Include in the global LOVD listing” setting, your LOVD installation will send us all the information we need to build these lists: the LOVD installation’s name, the URL, the available gene databases, the date of last update of the gene databases, the disease abbreviations and names of the diseases related to these genes and finally the names, institutes and email addresses of the curators of these gene databases. If you change this setting, please allow one day for the setting to take effect.

Security settings

In this section you can control some security settings of LOVD.

Allow submitter registration?
Enabling this setting allows submitters to create their own accounts in your LOVD installation. Having it enabled is the default setting, and this could not be disabled prior to LOVD 3.0-17.
Note, that submitters can never register when your LOVD installation is set to read-only because of an active announcement with the read-only setting enabled. See the Announcements section for more information.

Lock users after 3rd failed login?
With this setting enabled, submitters, curators and managers will be locked out of the system after they have provided the wrong password three times. A manager or, in the case of a locked manager, the database administrator needs to unlock the user’s account once it’s locked.
This does not affect the database administrator account.
Allow (locked) users to retrieve a new password?
If a submitter, curator, or manager has lost their password, enabling this “I forgot my password” feature allows them to receive a unlocking code in their email with which they can unlock their account and choose a new password.

Enable submitters to change data?
This setting allows submitters to make changes to data previously submitted by them or assigned to them.

Enable getting counts of hidden entries?
*Note: Not yet implemented!*
Enabling this feature allows the public to find the number of entries in the database (including hidden entries) matching one or more search terms on a specified set of columns. This feature will only mention the number of variant entries matched, without showing them.

Force SSL-only access to LOVD?
SSL is a secure protocol allowing for encryption of data sent between you and LOVD. When you record sensitive individual information in LOVD, you should enable this setting, as the individual information can otherwise be “sniffed” off the network. If you do not record sensitive information, enabling SSL is recommended.

Use data versioning of biological data?
*Note: Not yet implemented!*
Versioning allows you to see all previous versions of a certain data entry (individuals, variants, phenotype information, etc) and allows you to return the entry to a previous state. Please note that this feature requires quite a lot of space in the database. Disabling this feature later will not free any space, just prevent more space from being used.

4.2 System logs

System logs were introduced in LOVD 2.0 to allow authorized users to keep track of events in LOVD. This can be for security purposes, to follow each other’s progress when working together, or to track down errors in case something went wrong. The logs are initiated when LOVD is installed.

What is logged by LOVD
LOVD logs four types of events: authorization events and errors, general events, general errors, and installation and upgrade events. In case the event or error was triggered by an authorized user, the username of this user is also logged.

Authorization events and errors (“Auth”)
This list stores all successful and unsuccessful login attempts and password reset attempts. In all cases, the IP address from where the login or password reset attempts have been made, is also logged. For successful and unsuccessful login attempts the given username and the length of the given password is stored, so you can easily uncover brute force attempts where in general the attacker will try many random passwords. If a user can’t log in because of the IP restriction on the account, the current IP restriction is also stored in the log entry. Please note
that it is recommended to have LOVD lock user accounts after three failed login attempts. See subsection “Security settings” for more information.

**General events (“Event”)**

General events are logged to show a user’s activity, to allow collaborating users to track each other’s activities or to check what happened to a certain data entry over time.

**General errors (“Error”)**

Whenever an unrecoverable error occurs, LOVD logs the error and stops processing. Also some non-fatal errors are logged, such as when only part of the requested action could be completed. In principle, during any database event, an error can occur. This is then logged under the name of the event, so any of the event names used in the general event log can also be found in the error log.

**Installation and upgrade events (“Install”)**

LOVD creates a log entry on completion of the LOVD installation procedure. Also, after each upgrade, LOVD will add an entry stating from which version to which version it has been upgraded, and how many database queries have been used for the upgrade. Also errors during an LOVD upgrade, might they occur, will be logged here.

**Viewing and searching through the logs**

To browse the system logs, click the “View system logs” option of the “Setup” tab dropdown menu, or the “View, search and delete system logs” link in the setup area. The system log overview is fully searchable and sortable, allowing you to find log entries quickly.

The “Log” column shows the type of log every - Auth, Event, Error or Install. The “Date” column shows the date the log entry was created. The “User” column shows, if available, which user caused the log entry to be created. The “Event” column lists the type of event that occurred. Finally, the “Entry” column provides more information about the event that has occurred. For login attempts, it will contain the IP address of the user attempting to log in. For most event types, it will include the ID of the entry that was created, edited or deleted.

**Removing log entries**

To remove a single log entry, click on the red cross in the 6th column. To remove multiple log entries at once, first apply as many search terms on the log entries as needed to restrict the view to the entries you want to delete. Either manually select the checkboxes next to the entries you want deleted, or click the option button on the top left of the log overview, and select the “Select all ... entries” option, after which you can deselect log entries you do not want deleted. Then, open the option menu again and click the “Delete selected entries” option. Please note that deleting log entries can not be undone.
4.3 Announcements

For messages meant for visitors and users of your LOVD installation, you can create announcements. They will appear on the top of every page, can automatically appear and disappear on a requested date and time, and can optionally make LOVD read-only, disallowing submitter registrations and users lower than Managers to log into LOVD.

Creating a new announcement

To create a new announcement, click the “Create a new announcement, optionally making LOVD read-only” link in the Setup area, or click the “Create a new system announcement” link in the Setup tab drop-down menu.

Announcement details

Creating an announcement does not require many fields.

Announcement type

This defines the icon next to the announcement. An Information dialog has an information icon, a Question dialog has a question mark, a Warning dialog has an exclamation mark.

Announcement text

Type here the announcement itself. If you choose to make LOVD read-only as well, be sure to let users know from when they will be able to log in again.

Start date

Use this field to create an announcement before it should be online. The announcement will be active only after this date and time. If you leave this field empty, the current time will be selected, and the announcement will be shown immediately. To deactivate an announcement, put a date in the future, like 9999-12-31.

End date

Use this field to automatically let an announcement disappear at a certain date or time. If you leave this field empty, the end date will be set to 9999-12-31 23:59:59. To deactivate an announcement, put a date in the past here.

Make LOVD read-only?

Enabling this feature blocks logins and submitter registrations when this announcement is active. Only Managers and up will still be able to log into LOVD. Users of a lower level that are active when this announcement activates, will be logged out. A true read-only state is currently not enforced; Managers can still make changes to all data. Use this feature when the server is undergoing maintenance, for instance. If you want to make LOVD read-only, but only in the future, set a date in the future in “Start date”, and make sure you make an extra announcement that introduces the downtime.

After completing the form, click the “Create announcement” button. If the “Start date” field was empty or a day in the past, and the “End date” was empty or a date in the future, the announcement will be visible immediately. For Manager level users and up, announcements are clickable, and they link to their detailed view, making it easy for Managers to find active announcements.
4. LOVD Setup

4.4 Uninstalling LOVD

Only the database administrator can uninstall LOVD, provided the uninstall lock has not been set during installation of LOVD. The uninstall lock is set within the MySQL database and can only be removed by directly accessing MySQL. If you know how to handle MySQL, you can also uninstall LOVD through MySQL directly by removing all LOVD tables, but this is not recommended.

To uninstall LOVD and remove all data stored in LOVD, click on the “Uninstall LOVD” link in the setup area. This link is only available if the uninstall lock is disabled. You will need to fill in the database administrator’s password twice to complete the uninstallation process.

Once you have uninstalled LOVD, you have lost all variant and individual information. If you wish to store these data elsewhere, make sure you download all data first!

Please note that the LOVD uninstaller will not remove the LOVD files itself, just the database tables that it has created during install. Thus, to completely remove LOVD from the system, also remove the files from the server.
5. Authorized users

In LOVD 3.0, *authorized users* are everyone who registered for an account at the LOVD installation, including the database administrator who created their account during the installation process. The authorized users come in five levels; the database administrator of which there is only one, managers, curators, collaborators and submitters. The different levels of users are described in more detail below.

Registering for an account is necessary to be able to submit data or to become a gene curator. Directly after registering, you are allowed to submit individual, phenotype and variant data to the database, but the data needs to be checked and published by a *curator* or a *manager*. Curators and managers can submit and publish data in one step.

**User levels**

**Database administrator**

The database administrator is the first user within LOVD; this account is created during the installation of LOVD. After installation, the database administrator can create other users (managers, curators and submitters) in the system, if desired. These users can also register themselves, after which the database administrator can upgrade them.

The administrator has full access to all areas and settings of LOVD. The administrator is also the only user who can uninstall LOVD, provided the uninstall lock has not been set during LOVD installation.

When the “Forward messages to database admin” setting in the LOVD system settings is enabled, the database administrator will receive copies of submitter registrations and data submissions or edits.

**Manager**

A manager has access to almost everything the database administrator has access to. However, the manager is not able to create new managers or to edit existing managers. Also, a manager cannot uninstall LOVD even if the uninstall lock is not in place.

A manager can only be created by the administrator and often assists the administrator or takes over the customisation of LOVD for the curators.

**Curator**

The curator level user is actually a submitter, which has been given curatorship of one or more genes. Therefore, a curator only has access to the gene databases they are appointed to, and not to the system-wide setup area. However, also the database administrator or a manager can act as a curator by being appointed to certain genes, but in this case there is of course no restriction in access.

To assign a user to become curator, follow the steps explained in section “Assigning curators and collaborators” (available for managers and up).
Collaborator

The collaborator level user is a new type of user, introduced in LOVD 3.0. Collaborators are submitters with read access to non-public data on a certain given gene. Just like curators, these users must be appointed to genes by managers or the database administrator. See section “assigning curators and collaborators” for more information.

Submitter

Submitters have no special rights except to submit data to the database, which then needs to be checked and published by curators or managers. Depending on the LOVD system settings, submitters are also able to edit the data they have submitted previously.

Submitter accounts can also be created, edited or deleted by managers or the database administrator, although usually this should not be necessary. Curators do not require a separate submitter account; they can use their curator account to submit new data.

5.1 Registering a new account

Please note that you do NOT need to register to view the data available in the database. You only need an account for submitting new variant data.

Anyone can create a new submitter account. For collaborators, curators or managers, registering as a submitter is often the first step; their account can then be upgraded by a manager or the database administrator, respectively. The major difference between registration and direct creation of a new account by a manager or the database administrator, is that the registration can be performed by the public, and therefore it requires you to be logged out of LOVD. The registration form is protected using a reCAPTCHA module, which makes sure the form can only be used by humans, and not by spam bots.

To register a new submitter account, click the “Register as submitter” link on the top right corner of the screen, next to the “Log in” link. First, you are asked to provide an ORCID ID, if you have one. ORCID provides a persistent digital identifier that distinguishes you from every other researcher and, through integration in key research workflows such as manuscript and grant submission, supports automated linkages between you and your professional activities ensuring that your work is recognized. If you don’t have an ORCID ID yet, you can get one created by using the register link also given on the page. If you don’t want to register at ORCID at this time, click the “I don’t have an ORCID ID »” button to continue to the registration page. If you do have an ORCID ID, fill it in and click the “Continue »” button. Your ORCID ID will be verified, and the associated data is shown on the screen. If this data is correct, click “Yes, this is correct »” to continue to the registration page. As much information as possibly will be pre-filled for you. If the information shown is not correct, click “« No, this is not correct” so you can try and enter a correct ORCID ID or register without using one.
5. Authorized users

Registering a new account

User details
This part of the registration form contains the basic fields with which the user identifies themselves.

Name
Please enter your name in the way you want to be referenced to; usually in the format “<First name> <Last name>”.

Institute
The institute for which you work.

Department
If your institute consists of department, please enter the department for which you work.

Postal address
Please provide the full postal address of your institute.

Email address(es), one per line
Your account can hold multiple email addresses, which will all receive notifications sent by LOVD. This is useful if you’re sharing an account with a colleague, or when you would like to also get emails on a different account. You can enter as many email addresses as you like, one email address per line, no spaces, quotes or other special characters allowed.

Telephone
Optionally, provide your phone number where you can be reached.

Username
LOVD requires the username to be 4 to 20 characters long, and starting with a letter followed by letters, numbers, dots, underscores and dashes only.

Password
LOVD requires the password to be at least 4 characters long, containing at least one number or special character. Please choose your password wisely; don’t use the same password as your username and better not use words that are in a dictionary.

Password (confirm)
To make sure you didn’t make a typo while entering your password, please repeat it so we can check if the two fields are equal.

Referencing the lab
This part describes the reference that LOVD makes to your institute.

Country
Please select the country from the given list.

City
Even though you entered the city also in the address field above, please enter it here as well. It’s used for sorting purposes when creating a list of the institutes registered to the installation.

Reference
Your submissions will contain a reference to you in the format 'Country:City' by default. You may change this to your preferred reference here.
Security

To prevent unauthorized use of your account, we implemented a security feature that you can configure here.

Allowed IP address list

*Note: Please be extremely careful using this setting. Using this setting too strictly, can deny you access to LOVD, even if the correct credentials have been provided.*

An IP address is an address a computer is known by on the network. To help prevent others to try and guess your username/password combination, you can restrict access to your account to a number of IP addresses or ranges. For instance, to make sure your account can only be used when accessed from your institute or from your home.

Set to “*” to allow all IP addresses, use “-” to specify a range (like “192.168.0.1-20”) and use “;” to separate addresses or ranges (like “192.168.0.1;192.168.1.34”).

Registration authentication

To prevent automatic submission of the registration form, we implemented a reCAPTCHA module. In the provided text field, type the two words you see in the image. This will proof that you are a human and not a computer.

After completing the form, press the “Register” button. LOVD sends you a confirmation of your registered information through email. If the “Forward messages to database admin” setting in the LOVD system settings is enabled, the database administrator also receives a copy of your data.

5.2 Creating new user accounts

Besides the public registration form, managers can also create new users. This offers the advantage that the new user’s level can be set to the required level in the same step, and the manager does not need to log out first. You can create a new user by clicking on the “Create a new user account” option from the “Users” tab dropdown menu, or the “Create a new authorized user or submitter” link in the setup area. The process is largely equal to the submitter registration process, except that there is no reCAPTCHA test but instead you must enter your password for authorization, and the form has some additional fields, explained below.

User details

This section contains one additional field, compared to the registration page.

**Must change password at next logon**

If you want to force the user to pick a new password after logging in for the first time, select this checkbox. The user will not be able to work with LOVD until they have changed their password.
5. Authorized users

Security

This section contains more options when creating a new user account for a new user, compared with when registering your own account.

Level

Managers only see the option “Submitter” here. The database administrator may also select “Manager” here. Please note that to create a Curator account, you need to create a Submitter and then grant this user rights on the necessary genes. See section “Assigning curators and collaborators” for more information. For a complete description on all user levels, see the subsection “User levels”.

Locked

Users can be locked, which blocks access to LOVD even if they provide the correct username and password. This locking happens when too many times an invalid password was given while logging in with this username (depending on the LOVD system settings), or it can be done manually.

Other fields

This option is only available while creating a new user, it’s not shown on the submitter registration form.

Send email with account details to user

When enabled, the user’s details will be emailed to the email address(es) given, informing the user immediately about their account details. The submitter registration form does not have this checkbox, but this feature is always active for submitter registrations.

5.3 Editing user accounts

Every user in LOVD can edit their own account. Required level: Submitter

Editing other user’s accounts requires you to have a level higher than the user you are trying to edit. Managers can edit submitter accounts (including collaborators and curators). The database administrator can also edit manager accounts.

To edit your own account, click the “Your account” link on the top right-hand side of the screen, which leads directly to your account overview. Then, click the “Options” button and click “Update your registration”.

To edit someone else’s account, click the “Users” tab. Locate the user you wish to edit by searching or browsing through the given user listing. Clicking the entry will direct you to the user’s account details. If you have permission to edit this account, you’ll see an “Options” button. Click it and click “Edit user”.

The fields on the form are explained in the section “Registering a new account”. Please note that the “New password” fields are optional when editing an existing user account. You only fill them in when you want to change the password.

After you submit the form, LOVD redirects you automatically back to the user’s account details.
5.4 Locking users out of the system

A user is automatically locked if a wrong password is used three times while logging in with this username, provided this security setting is enabled (see section “LOVD system settings”), but this lock may also be placed onto, or removed from an account manually. Locking or unlocking a user account requires you to have a level higher than the user you are trying to lock or unlock. Proceed to the user’s account details page through the “Users” tab. Open the “Options” menu, and click the “Lock user” option. If the user is already locked, there will be an “Unlock user” option. The “Lock user” and “Unlock user” menu options are quick ways to deny or allow access to LOVD, although it can also be accomplished through the “Edit user” option.

Users who are logged in, can also be booted out of the system. If a user is logged in, the option “Force user log out” is added to the “Options” menu. This will destroy the user’s session, causing the user to log out. Locking the user essentially also causes the user to be logged out of the system the next time the user performs an action, although in that case it’s not possible for the user to log back in until the account is unlocked.

5.5 Sharing access with other users

An authorized user can share access to their data with other users. The other users can then see any non-public data entries owned by the sharing user, and if specified, they can also edit those entries. Additionally, all non-public fields of all entries owned by the sharing user, become available to this user, too.

A user can manage the people with whom they share their data from their account page. When logged in, there is a link named “Share access to your entries with other users” under the options menu on the user account page. When clicking this link, the user is presented with a page similar to that of Figure 5.1. This page shows in the bottom table the users to whom access is shared and whether they have edit permissions. The checkbox under “Allow edit” denotes whether the user has edit permissions on the shared data. One can remove users from this list by clicking the red cross next to their name, or add new users by finding and clicking their name in the top table. Finally, one can save changes to the bottom table by verifying their password and clicking “Save access permissions”.

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Figure 5.1: The “Sharing access” page, with a list of users one can select on top and a list of users previously selected on bottom.
6. Gene databases

Although in LOVD 3.0 you can enter data into the database without having genes in your installation, the genes still embody an important part of LOVD. Configuring genes in LOVD allows you to see the effects of variants on transcript level, and draws submitters with variant data to your installation.

Gene databases can also be created automatically for you, when you are importing VCF files in LOVD. However, this chapter deals with manually creating gene databases. You can create new genes through the LOVD setup area. Editing and deleting genes is done through the “Genes” tab. Curators assigned to genes can also edit these after they’ve been created, but can not delete genes. Emptying genes (removing all associated variant and patient data) and managing its variants is done through the configuration area. This chapter describes how to create, edit and delete gene databases in LOVD.

6.1 Creating a new gene database

Creating a new gene database in LOVD is largely automated. Click the “Create a new gene entry” option of the “Genes” tab dropdown menu, or the “Create a new gene database” link in the setup area. Fill in the HGNC ID or the gene symbol of the gene you want to create, and click “Continue »”. LOVD will check if the gene already exists in this LOVD installation, and if the gene symbol or HGNC ID is correct. For this, it contacts the HGNC website. If the gene symbol you used has been replaced by a new one, or if LOVD can find genes for which the given symbol is an alias, LOVD will report back the official symbol(s).

Unfortunately, due to some problems on the side of the NCBI, we are unable to handle variants on the mitochondrial genome at this time. NCBI does not handle the MT genes by their official names, and doesn’t allow for automatic retrieval of reference sequences for the MT genes. When the NCBI has fixed this problem, we will implement support for MT genes.

LOVD will show a progress bar while it contacts several webservice to retrieve gene information, reference sequence information and available transcript information. Usually retrieving this information takes about 1-3 seconds, but this depends on the internet connection between the server and the internet and the current load on the webservice which are contacted. When LOVD has retrieved all the information, you are forwarded to the data entry form where you can review the retrieved information and complete this gene’s settings.
General information

This part describes the bare essentials of the gene information. Therefore, most of the information in this section has been filled in for you, and some values are fixed to ensure the data is correct.

Full gene name
This information has been filled in for you, and can't be changed.

Official gene symbol
This information has been filled in for you, and can't be changed.

Chromosome
This information has been filled in for you, and can't be changed.

Chromosomal band
The chromosomal band on which the gene lies, is retrieved from the HGNC. You may modify it, if you wish.

Imprinting
If known, please fill in if this gene is imprinted (maternal or paternal) or not.

Date of creation
*Note: This field can only be filled in when creating a gene database and can never be edited afterwards!*
The database's date of creation is mentioned on the gene homepage. Today's date will automatically be stored if you leave this field empty, but in case you already had a gene database but you're moving it to this LOVD installation, you may want to fill in a different date.

Relation to diseases

This gene has been linked to these diseases
By linking the gene to disease(s), LOVD can help predicting next steps in data submission, and the link to the disease information will be shown on the gene homepage. If a disease you need is not present in this list, you can click the “configure more diseases” link below the selection list to create a new disease now. Afterwards, reload the gene form so you can select the new disease.
Alternatively, create the disease after the gene has been created.

Reference sequences

Collecting variants requires a proper reference sequence. Without a genomic and a transcript reference sequence the variants in this LOVD database cannot be interpreted properly or mapped to the genome. When properly configured, LOVD will add links to the Ensembl, NCBI and UCSC genome browsers from the gene homepage, generating a nice visual overview of the variants in the database.

Genomic reference sequence
Select the genomic reference sequence (NG or NC accession number). Only the references that are available to LOVD are shown. Checking for an available NG reference sequence is done at the NCBI. The NC reference sequence is always available, and depends on the chromosome on which the gene lies.
Transcript reference sequence(s)

LOVD retrieves a list of available transcripts from Mutalyzer. Mutalyzer currently only supports NCBI reference sequences. Please select which transcript you want to use to name the variants in this gene. If there are more transcripts, usually you would pick the transcript that encodes for the longest protein, or the transcript that is expressed in the tissue you focus your research on. If you want to add more transcripts later, or change which transcript(s) you want to use for this gene database, see the chapter Gene transcripts.

Links to information sources

Here you can add links to other resources that will be displayed on the gene’s LOVD gene homepage, such as Entrez Gene, OMIM, HGMD or GeneTests, but you can also add links to any website you want.

Homepage URL

If you have a separate homepage about this gene, you can specify the URL here. For the format: use the complete URL, including “http://” or “https://”.

External links

Here you can provide links to other resources on the internet that you would like to link to, such as other gene variant databases, or websites with more information about diagnostics. If you want to include more than one link, put every link on a new line in this field. If you do not want to include a title for this link, simply paste the full URL in this field, such as “http://DMD.LOVD.nl/”. If you do want to include a title, use the format “Description <URL>”, such as “Other LOVDs on this gene <http://DMD.LOVD.nl/>”.

HGNC ID

This information has been filled in for you, and can’t be changed.

Entrez Gene (Locuslink) ID

This information has been filled in for you, and can’t be changed.

OMIM Gene ID

This information has been filled in for you, and can’t be changed.

Provide link to HGMD

If you wish to include a link from the gene homepage to the gene’s page on the HGMD site, enable this checkbox.

Provide link to GeneCards

If you wish to include a link from the gene homepage to the gene’s page on the GeneCards site, enable this checkbox.

Provide link to GeneTests

If you wish to include a link from the gene homepage to the gene’s page on the GeneTests site, enable this checkbox.

This gene has a human-readable reference sequence

Although GenBank files are the official reference sequence, they are not very readable for humans. If you have a human-readable format of your reference sequence online, please select the type here; “Coding DNA” or “Genomic”.

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Human-readable reference sequence location

To create a human-readable reference sequence file, you can use our Reference Sequence Parser. It retrieves the reference sequence from Mutalyzer, and formats it into a human-readable format with good annotation of the upstream sequence, the exons, the introns, the downstream sequence, the predicted protein sequence and the alternative stop codons. Fill in the complete URL of the file, including “http://” or “https://”.

Customizations

LOVD allows you to customize the gene homepage by adding citation references, headers, footers, notes and a disclaimer. The header and footer will also be shown on the variant listings.

Citation reference(s)

Using our PubMed custom link, you can add links to PubMed articles to cite your database or that you want to point visitors to.

Include disclaimer

If you want a disclaimer added to the gene’s LOVD gene homepage, select your preferred option here.

Text for own disclaimer (HTML enabled)

Only in case you selected the “Use own disclaimer” in the field above, you can enter the text here. Text in this field will be ignored if you disabled the disclaimer or selected the standard LOVD disclaimer.

Page header (HTML enabled)

If you want a gene-specific header on all public gene-specific pages, please enter the text here.

Header aligned to

In case you defined a page header, here you can select how you want it aligned - left, center or right.

Page footer (HTML enabled)

If you want a gene-specific footer on all public gene-specific pages, please enter the text here.

Footer aligned to

In case you defined a page footer, here you can select how you want it aligned - left, center or right.

Notes for the LOVD gene homepage (HTML enabled)

Text entered here will appear in the General Information box on the gene’s homepage. In general, curators put additional information about the database here, such as how the data in the database was collected, grants supporting this work, or news.

Notes for the variant listings (HTML enabled)

Text entered here will appear below the gene’s variant listings.

Security settings

Using the following settings you can control some security settings of LOVD.

Allow public to download variant entries

This controls whether or not the public can also make downloads of your gene’s variants.
6. Gene databases

6.2 The gene homepage

LOVD automatically generates a gene homepage for every gene created within LOVD. The gene homepage is the entry point for visitors of a locus-specific database. In installations where LOVD is used for next-generation sequencing, the gene homepage serves more as a summary page of the gene-specific data in LOVD. Besides showing the basic information known about the gene in question (symbol, name, chromosome band, variant statistics, etc), curators also have quite a few options to customize the gene’s homepage.

If the gene you wish to view is already selected (its name is on the top of the page), simply click the “Genes” tab to open the gene’s homepage. Otherwise, move your mouse over the “Genes” tab, and select the “View all genes” menu option. Then, locate the gene you wish to edit by searching or browsing through the given genes listing. Clicking the entry will direct you to the gene’s homepage. Once on the gene’s homepage, the gene will be selected (its name will appear on the very top of the page), which allows faster navigation to information related to the gene of interest. For instance, when a gene is selected and you click the “Variants” tab, you are directed to the gene-specific variant overview. If no gene is selected, clicking on the same tab shows all variants in the database on the genome level. If you do not like this behavior, simply move your mouse over a tab, and select a different option from the dropdown menu.

6.3 Editing a gene database

Curators, appointed to a gene database, have the rights to edit its settings. Managers and up can edit all gene databases in the LOVD installation. To edit the gene’s settings, first proceed to the gene’s homepage.

Click on the “Options” menu button to open up the gene entry’s menu. If you do not see the “Options” menu button, you are not logged in or you do not have rights to edit this gene’s settings. In the opened menu, click “Edit gene information” to open up the gene’s edit form. The fields on the form are explained in the section “Creating a new gene database”. After you submit the form, LOVD redirects you automatically back to the gene’s homepage.

6.4 Deleting a gene database

Only managers and up can delete gene databases from LOVD. To delete the gene database in question, first proceed to the gene’s homepage.

Click on the “Options” menu button to open up the gene entry’s menu, and select the “Delete gene entry” option. To complete the removal, you need to fill in your password and submit the form.
Removing a gene can not be undone, you will need to create the gene database again. If in any doubt, first make a download of the database so that you can restore your data if needed. Please note that deleting a gene removes also its transcripts in the database and therefore all variant annotation on the gene in question, but it does not remove the genomic variants from the database. To remove these as well, you should first empty the gene database, which deletes all its variants and also deletes all other connected data such as individuals and phenotypes, as long as these are not connected to other genes as well.

6.5 Assigning curators and collaborators

Managers and up can assign curators and collaborators to gene databases, as well as revoke curator or collaborator rights. Directly after creating a new gene database, you are forwarded to the page that allows you to make these changes, but you can access that page at any time by using the “Add/remove curators/collaborators” link in the “Options” menu on the gene’s homepage.

![Authorize curators form](image)

**Figure 6.1:** The “Authorize curators” form, with on the top the users who are available to be selected, and on the bottom the selected curators and collaborators.

On the top of the page, you see a listing of all available users in the system, whichever level they have. Users that are already selected as curators or collaborators are shown in the bottom. To add a user as a curator or collaborator, search for the user in the top user list, and click the user entry. The user is then moved to the bottom.

In the bottom table, you see two checkboxes for each user. Also, you can sort the curators by dragging and dropping them in the list (see subsection Sorting curators). The first checkbox indicates whether or not they are allowed to edit the data in this gene’s database. Users allowed
to edit the data, are curators. Unchecking this checkbox turns this user in a collaborator for this gene; they can still see all public and unpublic data in this gene database, but they can’t edit it, like curators can. If the user is to be a curator, they will receive notifications of variant data submissions and submission updates. Collaborators do not receive such emails.

The second checkbox indicates whether or not the user’s name and email address is shown on the gene homepage and on the top of every page while this gene is selected. Curators can be hidden from the gene homepage and the page headers by unchecking this checkbox. Collaborators are always hidden, it is therefore not possible to change the second checkbox while the first checkbox is unchecked.

To remove an user as a curator or collaborator, click the red cross at the far right side of the table. You can only assign or remove curators if your user level is higher than that of the user you wish to assign/remove.

Please note that managers and the database administrator can never be made collaborators, since disallowing them to edit data contradicts their user level. They will always be allowed to edit all data in the system. Selecting them as curators doesn’t grant them additional access rights, but does enable the notification emails.

Also, please note that all genes in LOVD should have at least one visible curator. To be able to successfully submit the “Authorize curators” form, there needs to be at least one user with the “Allow edit” and “Shown” checkboxes checked.

### Sorting curators

![Figure 6.2](image)

**Figure 6.2:** The “Sort curators” form, allowing curators to resort the gene’s list of curator names, and show or hide curators from the gene homepage and the page headers.

Since the curator names are shown on the gene homepage and on the page headers, it is sometimes necessary to sort the curator names in a specific order. This can be done on the “Authorize curators” form (see figure 6.1 on page 36). Curators, who don’t have access to that form, use the “Sort/hide curator names” option from the “Options” dropdown menu on the gene homepage. This form greatly resembles the bottom part of the “Authorize curators” form, allowing to sort and show or hide the curator names (see figure 6.2 on page 37).

To resort the list of curators, move your mouse over the drag handle (see figure) on the left of the curator name, and click and drag the name up or down the list. Release the mouse button in the preferred location. Don’t forget to click the “Save” button to save the new sort order!
7. Gene transcripts

LOVD3 handles transcripts very differently from LOVD2. In LOVD3, configuring at least one transcript for a certain gene is necessary to be able to add variant data to the gene database. Genomic variants can always be added, but to record the effect of a variant on a certain gene, that gene must have at least one transcript defined. As a source for transcript information, LOVD relies on Mutalyzer, which gathers transcript information from different sources. Therefore, it is possible that new transcripts recently included in the NCBI database, are not yet available in LOVD because either the NCBI hasn’t rebuilt its mapping database yet, or Mutalyzer didn’t pick up the new mapping information yet. LOVD3 does not restrict you in the number of transcripts you can configure per gene, but note that for each transcript added, each variant gains more data fields that need to be filled in by the submitter, so make sure that the additional transcripts are actually needed.

7.1 Which transcript(s) should I pick?

In principle, we suggest using the predominant transcript, especially if only changes in this transcript cause a change in phenotype. Otherwise we recommend using the longest transcript (i.e., the transcript resulting in the longest protein). Using the predominant transcript makes confirmation of the variant effect on RNA level easier and usually the variant numbering on this transcript will be most familiar with other research groups. Using the longest transcript has the advantage of being able to annotate all exonic variants with a (predicted) effect on protein level.

As said, LOVD3 allows you to use more than one transcript per gene. Good examples that justify using multiple transcripts in LOVD, is when different transcripts are expressed in different tissues, when changes in different transcripts cause different phenotypes, or when different research groups focus on different transcripts (for example, when no transcript is available that contains all exons).

7.2 Creating new transcripts

The simplest way to create a gene transcript in LOVD, is to select it when creating a gene database. However, if you’d like to add more transcripts later, please proceed to the gene homepage. Click on the “Options” menu button to open up the gene entry’s menu. If you do not see the “Options” menu button, you are not logged in or you do not have rights to edit this gene’s settings. In the opened menu, click “Add transcript(s) to gene” to open up the gene’s edit form. Please wait while we contact Mutalyzer to provide a list of available transcripts. If you get the error message “No more transcripts available that have not been added yet!”, then all transcripts that are available to Mutalyzer have already been added to this gene.

Select the transcript(s) you would like to add, and click the button “Add transcript(s) to gene”. After the confirmation, you will be forwarded to the gene homepage. Beneath the header “Active transcripts” you can see which transcripts are now active.
7.3 Editing a transcript

Because LOVD standardizes the transcript information, there is very little information that is freely editable. At this moment, you are only allowed to assign or change Ensembl identifiers to the transcript(s) you use. To edit a certain transcript, proceed to the gene homepage, scroll down to “Active transcripts” and click the transcript in question. Alternatively, click the “Transcripts” tab from anywhere in LOVD. If you already had a gene database selected, you now see the transcripts active in this gene and you can click one to view its details. If you did not have a gene database selected, you now see all transcripts in the database. Use the filtering boxes to find the transcript you are looking for, and click it to view its details.

Click on the “Options” menu button to open up the transcript entry’s menu. If you do not see the “Options” menu button, you are not logged in or you do not have rights to edit this transcript’s settings. In the opened menu, click “Edit transcript information” to open up the transcript’s edit form.

General information

This part contains IDs to other sources, to aid in data exchange.

Transcript Ensembl ID

If this transcript is also known in Ensembl, enter its Ensembl ENST ID here.

Protein Ensembl ID

If this transcript’s protein product is also known in Ensembl, enter its Ensembl ENSP ID here.

Protein Uniprot ID

If this transcript’s protein product has an Uniprot ID, enter it here.

7.4 Deleting a transcript

Deleting a transcript removes all information related to the transcript in question from all variants in the database. Just like when deleting a gene, this does not remove the genomic variants from the database! From the transcript’s detailed view, click on the “Options” menu button to open up the transcript’s menu, and select the “Delete transcript entry” option. On the next page, confirm by typing your password in the input box and submit the form to complete the removal.
8. Custom columns

To make sure LOVD can be used by a great variety of users who all focus on storing different data, LOVD allows great flexibility in the form of custom columns; data fields that can be personalized by the curators, or even completely newly defined. All standard LOVD columns can be edited and removed, except for the HGVS recommended columns, which can only be edited but not removed. Besides this default set of selected columns which are added by default, LOVD also comes with a set of columns that have not been added to the data tables yet, but can be added with a click of a button. Additional columns can be created to suit your every need.

8.1 Custom column categories

In LOVD3, custom columns come in five different categories.

Individual
Information on the individual, not related to disease, not changing over time, such as date of birth, geographic or ethnic origin.

Phenotype
Information on the phenotype, related to disease, possibly changing over time, such as blood pressure, body length or date seen by specialist.

Screening
Information on the detection of new variants, such as detection technique, laboratory conditions or type of sample taken from the individual.

Variant On Genome
Information on the variant(s) found, in general or on the genomic level, such as restriction site change, DNA change on genomic level, or reference describing the variant.

Variant On Transcript
Information on the variant(s) found, specific for the transcript level, such as predicted effect on protein level, DNA change on mRNA level, or exon number.

Of these five different categories, two are shared: they can be turned on or off by curators for specific diseases or genes, their so-called parent objects. Some of these columns’ settings can also be changed per parent object they are enabled in. The two shared custom column types are “Phenotype”, configurable per disease, and “Variant On Transcript”, configurable per gene.

The other three categories are system-wide; meaning they are either always on (enabled), or always off (disabled). These column are referred to as non-shared columns, and can only be enabled, disabled and managed by Manager level users, or the Database administrator. The three non-shared custom column types are “Individual”, “Screening” and “Variant On Genome”.

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8. Custom columns

8.2 Creating a new custom column

By creating a new custom column, you can define what kind of information you want to store in the database. Please note that defining this type of information, does not automatically make LOVD store this information. You will need to enable the column after defining it, so it actually gets added to the data entry form and show up in the data tables.

To create a new custom column, click on the “Create new custom data column” option from the “Setup” tab dropdown menu, or the “Create new custom data column” link in the setup area. Firstly, you will need to choose which type of column you would like to create. For more information on the different column types, see the previous section. Please note that you can never change the type of a column once it has been chosen. Select the type of column you would like to create, by clicking its description. You are then forwarded to the entry form specific for this type of custom column.

Column name and descriptions

Here the description of the new column is defined; its displayed name and the explanation of its contents to the users.

Column ID

Choose a simple unique name for your column, consisting of only letters, numbers and underscores. Subcategories must be divided by a slash (/), such as “Geographic_origin/Country”, “Blood_pressure/Systolic”, “Protocol/Date_updated”, “Frequency/dbSNP” or “Protein/Codon”.

Column heading

This will appear above the column in data tables and on the legend. In principle, this is free text, but try to keep it short, because a long name will increase the width of the column.

Description on short legend (HTML enabled)

For shared columns, this value can be changed separately for each parent object this column has been added to.

Describe the meaning of the column in short. This will appear when a user moves their mouse over the column header in the data tables.

Description on full legend (HTML enabled)

For shared columns, this value can be changed separately for each parent object this column has been added to.

Describe the meaning of the column. You can be very elaborate here. This text will be shown on the full legend of the data table.

Data and form settings

Here you define what the column looks like on the data entry form, and what kind of data it will store. There are two ways to define the data type of your custom column. The recommended way is to use the data type wizard, which will fill in the MySQL and the form type values. Only if you really know what you’re doing, you can edit the MySQL data type and the form type directly. Its formats are out of the scope of this manual, and therefore not explained here.
Using the data type wizard

To start the data type wizard, click the “Start data type wizard” button. The wizard will open in a new window.

Firstly, you need to select what type of data entry field you want to create for your custom column. In the “Basic form style” field, choose from one of these options:

**Text/numeric input field**
A standard input field allowing textual or numerical free text input.

**Integer input field**
A standard input field, visually similar to the text/numeric input field, but allowing only whole numbers to be filled in.

**Decimal input field**
A standard input field, visually similar to the text/numeric input field, but allowing only numbers with a configurable number of decimals.

**Large multi-row textual input field**
A large text field with multiple rows, allowing large amounts of text to be filled in.

**Drop down list (one option selected)**
A drop down list with a set of options of which the user must choose one value.

**Selection list (multiple options selected)**
A selection list where the user can select one or more options or ranges.

**Date input field**
A standard input field, visually similar to the text/numeric input field, but allowing only dates (year, month, day), optionally also storing time (hours, minutes, seconds). The input format is YYYY-MM-DD or YYYY-MM-DD HH:MM:SS.

**On/off checkbox**
A simple checkbox allowing only a yes/no answer.

Select the wanted data entry type and click “Next »”.

The next page shows you some additional options that depend on the type of field you’ve chosen.

**Column name on form** (all field types)
The name the column will have on the data entry form. This is usually the same as the “Column heading” field.

**Help text** (all field types)
If you think the data field needs clarification for users, especially those who don’t know this field well, add such clarification here. By filling in this field, LOVD will add a question mark icon next to the field on the data entry form. When the user moves their mouse over the question mark icon, the text entered here is shown. Since the text is shown as a highlight, it does not disrupt the structure of the data entry form, so feel free to type lengthy texts.
8. Custom columns

Creating a new custom column

Notes on form (all field types, HTML enabled)

For shared columns, this value can be changed separately for each parent object this column has been added to.

If you think the data field needs clarification for all users, add such clarification here. The text given here will be shown directly underneath the data entry field. Therefore, keep it short, since long texts will make the data entry form very long.

Width on form (characters) (all input fields and text area)

The input field width on the form, in number of characters.

Height on form (lines) (text area and multiple selection list)

The height of your field on the data entry form, in lines. For multiple selection lists, this implies the number of options visible at the same time.

Maximum input length (characters) (text and integer input field types)

The maximum number of characters allowed the be filled in. If a user tries to enter more characters in your field, an error is issued. For the integer input type, this implies the maximum value that can be filled in.

Number of digits before the decimal point (decimal field only)

The maximum number of digits allowed before the decimal point. This practically limits the maximum value of the field.

Number of digits following the decimal point (decimal field only)

The maximum number of digits allowed at the right of the decimal point. This limits the maximum precision of values in the field.

Regular expression pattern (text input only)

For advanced users only. You can enter a full regular expression pattern (PHP’s Perl-compatible regular expression syntax), including ‘/’ delimiters and possible modifiers. Using this, you can force a certain format for the input. Make sure it’s valid, otherwise you risk getting all this column’s data input rejected.

Allow only positive values (integer and decimal fields)

Select this to allow only positive numbers to be entered.

Provide “– select –” option (drop down list only)

This will add an option named “– select –” to the list of options, that will be selected by default and will be regarded as an empty value. If this option is not set, the first option of the list will be the default selected option.

Provide “select all” link (multiple selection list only)

This will add a link next to the selection list that allows the user to instantly select all available options.

List of possible options (both list types)

For shared columns, this value can be changed separately for each parent object this column has been added to.

Enter the options available for this field, one options per line. If you want to use abbreviations, use: Abbreviation = Long name, like “DMD = Duchenne Muscular Dystrophy”. In that case the user will see the long name in the selection list, but the abbreviation will be stored in the database.

Also store time? (date field only)

Select this to also store time (hours, minutes, seconds) in this date field.
8. Custom columns

Creating a new custom column

**Default value** (all input fields except text area)

A default value that will appear in the field when creating a new data entry.

Click the “Finish” button to have the data type wizard fill in the “MySQL data type” and “Form type” fields for you, based on the information you provided.

**Column settings**

Here you can change some of the column’s settings. For shared columns (see section “Custom column categories”), most of these settings can also be changed per parent object where this column has been added to.

**Include this column for newly configured genes/diseases** (shared columns only)

When this is selected, new genes (for VariantOnTranscript columns) or diseases (for Phenotype columns) will have this custom column enabled by default. When you leave this off, the curators of new genes or diseases will have to enable the column themselves, should they want to use it.

**Column display width in pixels**

For shared columns, this value can be changed separately for each parent object this column has been added to.

Here you can define how wide the column should be in data listings. This is measured in pixels. Since the width depends on the screen’s resolution, a hint is provided how wide the currently selected width is on the current screen.

**Mandatory field**

For shared columns, this value can be changed separately for each parent object this column has been added to.

With this option selected, users can’t leave this column empty.

**Show contents to public**

For shared columns, this value can be changed separately for each parent object this column has been added to.

This controls whether or not users with no specific authorization on the data entry can see the contents of this column.

**Show field on submission form**

For shared columns, this value can be changed separately for each parent object this column has been added to.

If you don’t want submitters to fill in this column, but reserve the use for curators, disable this checkbox. Please note that unsetting this will not hide the contents of the column from the public, it merely stops them from entering data in it.

**Link settings**

Here you can select which custom links will be active in this custom column.

**Active custom links**

Select any number of custom links, or leave all unselected if you do not wish to use custom links in this field. Note that the use of custom links only makes sense for fields that accept text input.

Enter your password at the bottom of the form and submit it. Your custom column is now created, and you can choose to enable the column if you wish. Also, you might want to check the column order, since newly created custom columns are appended to be the last column.
8.3 Enabling custom columns

Directly after creating a new custom column, or at any later time, you can enable a custom column. It must already exist in the database, otherwise you first need to create it. Non-shared custom columns (Individual, Screening and Variant On Genome categories) are enabled system-wide, which requires manager access. Shared custom columns (Phenotype and Variant On Transcript categories) can be enabled by parent-object (disease or gene, respectively) and require curator access.

Enabling non-shared custom columns

The non-shared, system-wide custom column types are Individual, Screening and Variant On Genome. To enable a non-shared custom column, you should start from the list of currently available columns. You can get there in two ways:

- Depending on which type of column you wish to enable, click the “Enable more custom columns” link in the “Individuals” or “Screenings” menu tabs’ dropdown menu, or the “Enable more genomic custom columns” link from the “Variants” menu tab dropdown menu.

- Click the “Browse all custom data columns” link of the setup menu tab dropdown menu, or from the setup area, click the “Browse all custom data columns already available to enable or disable them, or view or edit their settings” link listed under “Custom data columns”. This is the list of all custom columns. To filter this list for category, click the options wheel icon on the top left hand side of the columns listing to open up the menu, and select the category you are interested in. You can also filter for category by typing (part of) the name in the “Category” column filter box, and press the “Enter” key. To further filter the list for columns not yet added to the data table, fill in a zero (‘0’) in the “Active” column filter box and press the “Enter” key.

From the column overview, click the column you wish to enable.

In the detailed view of the custom column, you can review all the column’s settings to verify that it suits your needs. Click the “Options” menu button to open up the column’s menu, and click on “Enable column”. Please note that when your database already contains a lot of information, enabling a new non-shared column may take some time. If this is the case, LOVD will display a warning. Type in your password in the “Enter your password for authorization” field for verification of your authorization, and submit the form. The column is now enabled and visible in the data entry forms and the data listings and detailed views. You may want to review or change the order of the custom columns now.

Enabling shared custom columns

The shared custom column types are Phenotype and Variant On Transcript. Before enabling more shared columns, you might first want to check out the default order in which they will appear. To change the default order for shared custom columns, see subsection “Changing the system-wide custom column order”.

To enable a shared custom column, you should start from the list of currently available columns. You can get there in two ways:
8. Custom columns

8.4 Editing custom column settings

Shared columns have some settings, mostly related to the display of the column, which can be set per parent object (a certain disease or gene). We will refer to these settings as object-specific settings. All other settings are system-wide and can only be performed by a manager or higher. Non-shared columns only have system-wide settings and can not be configured per disease or gene.

Editing system-wide custom column settings

Both shared and non-shared columns have system-wide settings that can be changed by a manager or higher. However, the way these columns respond to changes to the system-wide settings are not the same. For non-shared columns, each change to the system-wide settings will be effective immediately. For shared columns, only the settings that are not object-specific (see above) are effective immediately. All changes to object-specific settings however, come to serve as new default values for when the custom column in question is activated for new genes or diseases. Already enabled shared custom
columns will not have their object-specific settings reset, unless they are explicitly reset to the (new) default settings (described in the paragraph below).

To edit a custom column’s system-wide settings, you should start from the list of currently available columns. Click the “Browse all custom data columns” link of the setup menu tab dropdown menu, or from the setup area, click the “Browse all custom data columns already available to enable or disable them, or view or edit their settings” link listed under “Custom data columns”. This is the list of all custom columns. To filter this list for category, click the options wheel icon on the top left hand side of the columns listing to open up the menu, and select the category you are interested in. You can also filter for category by typing (part of) the name in the “Category” column filter box, and press the “Enter” key. Now, click the column which settings you wish to edit. Click the “Options” menu button to open up the column’s menu, and click on “Edit custom data column settings”.

The form to edit the system-wide settings is equal to the form used to create a new custom column, with two exceptions. The “Column ID” field is not present here, since you can not change a column’s ID once you created it. Also, for shared columns, at the bottom of the form a checkbox is present labeled “Apply changes to all genes/diseases where this column is active”. Checking this checkbox effectively resets all enabled instances of the column and applies all changes you make on this form to them, overwriting possible modifications of object-specific settings. All other fields on this form are explained in the section “Creating a new custom column”.

**Editing object-specific (shared) custom column settings**

The object-specific settings, editable per gene or disease, are (also see section “Creating a new custom column”):
- Description on short legend
- Description on full legend
- Notes on form
- List of possible options (for selection list type columns)
- Column display width in pixels
- Mandatory field
- Show contents to public
- Show field on submission form

To change these settings, first you need to proceed to the detailed view of the disease or gene in question. For Phenotype columns, move your mouse over the “Diseases” menu tab, and click the “View all diseases” link from the dropdown menu. For Variant On Transcript columns, move your mouse over the “Genes” menu tab, and click the “View all genes” link from the dropdown menu. Then, select the parent object you would like to change the object-specific settings for.

From the parent object’s detailed view, open the “Options” menu and click the “View enabled phenotype columns” or “View enabled variant columns” link. From the list of enabled columns, click the one you wish to edit the settings for. Open up the “Options” menu and then click “Edit settings for this disease only” for Phenotype columns, or “Edit settings for this gene only” for Variant On Transcript columns. For explanation of the available fields, please see the section “Creating a new custom column”. After submitting the form, the changes are effective immediately, but applied to the chosen parent object only.
8. Custom columns  

8.5 Changing the order of custom columns

Custom columns are shown in a predefined, but adaptable order. This order determines how custom columns are shown in the data tables, in the download files, and on data entry forms. It is therefore important that the columns are sorted in a sensible order.

Just like the custom column settings, shared columns can have an object-specific order, different from the default order. Similarly, the order of non-shared columns is system-wide and needs to be configured by a manager or up.

Please note that columns from different categories can not be mixed. Some data listings show overviews of different types of data joined together, such as variants on genomic level joined with the variant’s effect on transcript level. In this case, the columns representing genomic data, such as chromosome, DNA description on genomic level and the allele field, can not be mixed with columns representing data related to the change on the transcript, such as exon number, DNA description on the transcript level or the effect on protein level. Only the order in which each category’s columns are shown, can be changed.

Changing the system-wide custom column order

For non-shared columns, changing the system-wide custom column order will effectively change the current order in which they are shown in data listings and data entry forms. For shared columns, changing the system-wide order will merely change the default order in which the columns appear when new parent objects (diseases, genes) are created. It also influences where a newly enabled column will show up amidst the columns already active for the object it just got added to. The order of already enabled shared custom columns will not be affected.

To change the custom column’s system-wide order, you should start from the list of currently available columns. Click the “Browse all custom data columns” link of the setup menu tab dropdown menu, or from the setup area, click the “Browse all custom data columns already available to enable or disable them, or view or edit their settings” link listed under “Custom data columns”. This is the list of all custom columns. To filter this list for category, click the options wheel icon on the top left hand side of the columns listing to open up the menu, and select the category you are interested in. Now, click the options wheel icon again, and click the “Change order of columns” link. Alternatively, first select one of the columns from the selected category, and on the detailed view open the “Options” menu and click the “Re-order all «category» columns” link.

You now see all custom columns in this category, both active and not active. On the left of each column’s header you see a handle (see figure) to drag the column up or down. To change the order in which the columns are shown, move your mouse over the drag handle, and click and drag the name up or down the list. Release the mouse button where you wish to drop the column. Don’t forget to click the “Save” button to save the new sort order!
8. Custom columns

Disabling custom columns

8.6 Disabling custom columns

Custom columns that you no longer wish to use, can in most circumstances be removed by disabling the column. Exceptions in this case are HGVS standard columns. HGVS standard custom columns are columns that are part of the mandatory minimal data set as defined by the HGVS for Locus-Specific DataBases (LSDBs). This is to preserve the minimal quality of the data within the database, and LOVD will not allow you to remove these columns. Disabling non-shared custom columns is a system-wide change, which requires manager access. Disabling shared custom columns can be done per parent-object and requires curator access.

Disabling non-shared custom columns

To disable a non-shared custom column, you should start from the list of currently active columns. You can get there in two ways:

- Depending on which type of column you wish to disable, click the “View active custom columns” link in the “Individuals” or “Screenings” menu tabs’ dropdown menu, or the “View active genomic custom columns” link from the “Variants” menu tab dropdown menu.

- Click the “Browse all custom data columns” link of the setup menu tab dropdown menu, or from the setup area, click the “Browse all custom data columns already available to enable or disable them, or view or edit their settings” link listed under “Custom data
columns”. This is the list of all custom columns. To filter this list for category, click the options wheel icon on the top left hand side of the columns listing to open up the menu, and select the category you are interested in. You can also filter for category by typing (part of) the name in the “Category” column filter box, and press the “Enter” key. To further filter the list for columns currently enabled, fill in a one (‘1’) in the “Active” column filter box and press the “Enter” key.

From the column overview, click the column you wish to disable.

Click the “Options” menu button to open up the column’s menu, and click on “Disable column and remove values”. Please note that when your database already contains a lot of information, disabling a non-shared column may take some time. If this is the case, LOVD will display a warning. Type in your password in the “Enter your password for authorization” field for verification of your authorization, and submit the form. The column is now disabled and removed from the data entry forms and the data listings and detailed views.

If the “Disable column and remove values” link is not enabled, i.e. you see it, but you can’t click it, the column is an HGVS standard column, and it can not be removed.

Disabling shared custom columns

Required level: Curator
Available from: LOVD 3.0 Build 01

Shared custom columns can be disabled for one parent object only, or from multiple parent objects at the same time.

To disable the given column for one parent object only, proceed to the detailed view of this parent object (disease or gene). Then, open the “Options” menu button and click the “View enabled phenotype columns” or “View enabled variant columns” link, depending on the type of column you wish to disable. Click the column, then from its “Options” menu button, select “Remove column from this disease” or “Remove column from this gene”, depending on the type of column. Please note that if this shared custom column will no longer be active for any parent object and your database already contains a lot of information, disabling this column may take some time. If this is the case, LOVD will display a warning. Type in your password in the “Enter your password for authorization” field for verification of your authorization, and submit the form.

If the “Remove column” link is not enabled, i.e. you see it, but you can’t click it, the column is an HGVS standard column, and it can not be removed.

To disable the given column for multiple parent objects at the same time, start with one of the following steps.

- From any disease or gene that you have authorization of, open the “Options” menu button and click the “View all available phenotype columns” or “View all available variant columns” link, depending on which type of column you wish to disable.

- If you’re a manager or higher, click the “Browse all custom data columns” link of the setup menu tab dropdown menu, or from the setup area, click the “Browse all custom data columns already available to enable or disable them, or view or edit their settings” link listed under “Custom data columns”. This is the list of all custom columns. To filter this list for category, click the options wheel icon on the top left hand side of the columns listing to open up the menu, and select the category you are interested in. You can also filter for category by typing (part of) the name in the “Category” column filter box, and press the “Enter” key.
To further filter the list for columns currently enabled, fill in a one (‘1’) in the “Active” column filter box and press the “Enter” key. Please note that in this custom column overview, the “Active” field simply states whether or not this column has been added to the data table, i.e. is active for any parent object.

From the column overview, click the column you wish to disable.

In the detailed view of the custom column, open its “Options” menu button, then select “Disable column”. Please note that if this shared custom column will no longer be active for any parent object and your database already contains a lot of information, disabling this column may take some time. If this is the case, LOVD will display a warning. From the “Remove this column from” selection list, select the parent objects (diseases or genes) you wish to remove this column from. For help on how to use this type of selection list, move your mouse over the blue help icon. If the parent object you are looking for is not shown in this list, you have no authorization on the object to change its column’s settings. Finally, type in your password in the “Enter your password for authorization” field for verification of your authorization, and submit the form. The column is now disabled and removed from the data entry forms and the data listings and detailed views.

If the “Disable column” link is not enabled, i.e. you see it, but you can’t click it, the column is an HGVS standard column, and it can not be removed.

### 8.7 Deleting custom columns

Custom columns, which are created by users (managers or the database administrator), can also be deleted permanently from the system, for instance if they were for testing purposes only or if they are no longer needed. First, you need to make sure the column is no longer active. See section “Disabling custom columns” on how to inactivate a column.

First, proceed to the column’s detailed view; access the custom data columns listing through the setup menu tab dropdown menu or setup area, find the column you wish to delete, and click on its entry. Then, from its “Options” menu button, select “Delete column”. Type in your password in the “Enter your password for authorization” field for verification of your authorization, and submit the form.

If the “Delete column” link is not enabled, i.e. you see it, but you can’t click it, the column is either still active (check the ‘Active in LOVD?’ field), it’s an HGVS standard column (check the ‘HGVS required column’ field), or it is created by LOVD itself (check the ‘Created by’ field). In any of these cases, the column can not be removed.

### 8.8 Downloading custom column data to text files

LOVD supports exporting the custom data columns, including the columns’ system-wide settings, to the LOVD import format. This file can then be used as a backup or to share the custom columns specifications with other LOVD3 installations, by (re)importing the file into LOVD (from LOVD 3.0-05). Please note that from LOVD 3.0-05, the custom column data is also included in the full LOVD download, available from the setup area.
To download all custom column specifications, click the “Download all LOVD custom columns” link in the setup menu tab dropdown menu, or from the setup area, click the “Download all LOVD custom columns in the LOVD import format” link listed under “Custom data columns”. We do not recommend you to edit this file.

For more information on the format, please see section “The LOVD3 import format”.

**8.9 Importing custom column data from text files**

Custom columns downloaded in the LOVD import format can be reimported into LOVD. LOVD will only import the custom columns from the file which are not already in the system. LOVD does not compare the column’s settings in the file versus what is in the database, LOVD just checks if the column IDs are present in the database or not.

Please see section “Importing data into LOVD” on how to import data into LOVD.
9. Adding data to LOVD

All registered users can submit or add data to LOVD. A complete submission contains information on an individual, disease and phenotype information (optional), variant screening(s), and at least one found variant. You can add more information to your submission at any later time, like new variants, phenotype information or a new screening. If you are a submitter level user, all new data submissions have to be published by a curator before the data is visible to the public. When finishing a submission, the curator(s) will receive an email notification, inviting them to publish the data.

This chapter describes adding data to LOVD using the web forms. If you are a curator or up and you wish to add data to LOVD using a tab-delimited file upload, please see chapter “Downloading & importing data”. If you’re a programmer and you would like to submit data in JSON format directly from your application using the LOVD3 submission API, please see chapter “The LOVD APIs”.

9.1 Starting a submission by creating an individual

You can submit new data by clicking the “Submit” tab, or by selecting the option “Create a new data submission” from the dropdown menu of the tabs “Individuals”, “Screenings” or “Variants”. Depending on your authorisation level, you have different options for submitting data. As an submitter level user, your submission always starts directly with creating an individual to whom you can link phenotype or variant screening information. If you are registered as a curator or higher, choose “I want to submit information on individuals” to enter individual information. As a curator or higher, you can also choose to submit only summary variant data, although we do not recommend this. Please see section “Submitting summary variant data” for details.

In the data entry form for data on the individual, most fields are optional in a standard LOVD installation. You can check the help icon next to the form fields to see details about the fields and suggestions on how to use the fields. After completing the form, click “Create individual information entry”. When successful, a message is displayed several seconds: “Successfully created the individual information entry!”. Hereafter, you are redirected to the menu as displayed in figure 9.1, where you can choose how to proceed.
Adding phenotype information to the individual

When you created an individual, and this individual has been diagnosed with one or more diseases, or when this individual is marked as healthy/control, it is possible to add extra phenotype details to this individual. Select “I want to add phenotype information to this individual” (figure 9.1), fill all required fields and click “Create phenotype information entry”.

The option “I want to add phenotype information to this individual” is disabled when no relation to diseases has been selected during the creation of this individual. If you wish to add phenotype information while this option is disabled, you have to edit the individual, see section “Adding phenotype information to an individual” on how to proceed.

Adding a variant screening to the individual

By selecting “I want to add a variant screening to this individual” you can add a screening to the individual. For “Detection template”, “Technique(s) used” and “Genes screened”, more than one item can be selected. Complete the form, and when ready click “Create screening information entry”. When the entry is successful, a message is displayed for several seconds: “Successfully created the screening entry!”. Hereafter you are redirected to a menu as displayed in figure 9.2, where you can choose how to proceed.

Adding a variant to a screening

After adding a screening, you can add a variant to this screening. It is possible to add more than one variant to a screening. A submission can only be finished when at least one variant has been added to the submission.

Confirming variants using a screening

When different variant screenings have been applied to the individual, for instance to confirm a variant found using a different method, the screenings can be stored separately in the database.
Multiple screenings can be linked to the same variant, or have their own sets of variants. The option to confirm variants using a screening is only available when at least two screenings are added to the individual and variants have already been submitted. After choosing this option, you are shown a list of variants already linked to other screenings of the individual, but not to the currently selected screening (see figure 9.4). Click the checkmarks left of the variants that you would like to mark as confirmed, and click “Save variant list”.

Adding a variant located within a gene

After choosing this option, you are shown a list of genes (see figure 9.5). If certain genes have been selected while creating the screening, these are pre-selected. You can remove or edit the pre-selection by removing or editing the configured filter in the “Symbol” column. Then click the gene for which you want add a variant to proceed to the variant entry form. You can check the help icon next to the form fields to see details about the fields and suggestions on how to use the fields. Complete the form, and when ready, click the “Create variant entry”.

Figure 9.2: After adding a screening, you can add a variant to this screening. “Back to the individual” will leave this screening unfinished, and will return you to the choice to add phenotype information or another screening.

Figure 9.3: Creating a new variant entry. Note: Only managers and up have the option “I want to upload a file with genomic variant data” available.
Adding data to LOVD

Starting a submission by creating an individual

Figure 9.4: Select a variant to confirm. Shown are the variants already linked to other screenings of the individual, but not to the currently selected screening.

button. If successful, you are redirected to the options to submit more variants, or return to the individual.

Figure 9.5: Click the gene for which you want to submit a variant.

Adding a variant only described on genomic level

You can create a variant which is only described on genomic level. The data entry form is similar to the form for adding a variant within a gene, but lacks the transcript-specific fields. Check the help icons next to the form fields to see details about the fields and suggestions on how to use the fields. Complete the form, and when ready, click the “Create variant entry” button. If successful, you are redirected to the options to submit more variants, or return to the individual.

Uploading file with genomic variant data

If you are a manager or higher, you can upload a file with genomic variant data. This option is not displayed for users with levels lower than manager. For uploading genomic variant data
files, the Variant Call Format (VCF) version 4.0 or higher is supported.

### Adding a variant to the individual

The option “Add a variant to this individual” as displayed in figure 9.1 is only available when you already created a screening, and you selected “Back to the individual” after creating a screening.

When you want to add a variant to the individual, using a screening you previously created, you have to select the appropriate screening. Click the “Add a variant to this individual” option, then click the screening you wish to add another variant to, and you will be redirected to the menu where you can submit a new variant. See section “Adding a variant to a screening”.

### Finishing the submission

The option “I want to finish this submission” is enabled for submitters when they completed a submission, i.e., an individual, screening and at least one variant have been added. Finishing a submission will send an email to the curator(s) of the gene(s) affected by the variant(s) submitted, or the manager(s), for variants described only on genomic level. They are thereby requested to publish the newly submitted data. You as submitter of the data, will receive a copy of this email. As long as the data is not yet published, the data is invisible for other users. You will however be able to see the data previously submitted by you, at all times.

### 9.2 Continuing an unfinished submission

If, for whatever reason, a data submission has not been finished, it is stored in the list of unfinished submissions. This also means that the curator(s) did not get an email with the request to publish the newly submitted data. You can continue an unfinished submission at any time, adding any missing information, and finishing the submission to alert the curator(s) of the new data. To proceed with an unfinished submission, select “Unfinished submissions” in the top right corner of the screen. You will see an overview of data submissions which are not yet finished. Select any submission to proceed. The normal submission process will continue where it was left, for instructions see the appropriate section in this chapter.

### 9.3 Submitting summary variant data

If you are a curator or higher, you can choose to submit only summary variant data. This means that no individual, phenotype or screening data is submitted, only variant data. Although you can submit summary variant data, it is highly discouraged because individual and phenotype data make the variant data much more valuable. An example of a valid reason to submit summary variant data is for storing variant frequency data of large populations.
9.4 Adding data to already finished submissions

It is still possible for the submitter to add data to an individual or a screening when a submitter finished the submission, regardless of whether the curator published the data or not.

Adding a variant to a screening

To add a variant to an existing screening, click on the “Screenings” tab and select a screening for which you are authorized. Click on the “Options” menu button to open up the screening entry’s menu. If you do not see the “Options” menu button, you are not logged in or you do not have rights to edit this screening. In the opened menu, click “Add variant to screening”. From here you can follow the steps described in the section “Adding a variant to a screening”. After you submitted the variant data, LOVD sends an email to the curator(s) and redirects you to the variant overview.

Adding screening to individual

To add a screening to an existing individual, click on the “Individuals” tab and select an individual for which you are authorized. Click on the “Options” menu button to open up the individual entry’s menu. If you do not see the “Options” menu button, you are not logged in or you do not have rights to edit this individual. In the opened menu, click “Add screening to individual” to open up the screening data entry form. You are now in the submission process described in section “Adding a variant screening to the individual”. The option “Back to individual” is not available when you submit a screening to an existing individual. You can only finish this submission, if you also add at least one new variant to this new screening, or select at least one previously submitted variant as confirmed by this new screening. After finishing the submission, LOVD sends an email to the curator(s) requesting the publication of the submission.

Adding phenotype information to an individual

To add phenotype information to an existing individual, click on the “Individuals” tab and select an individual for which you are authorized. Click on the “Options” menu button to open up the individual entry’s menu. If you do not see the “Options” menu button, you are not logged in or you do not have rights to edit this individual. In the opened menu, click “Add phenotype information to individual” to open up the phenotype data entry form, or, in the case this individual has been diagnosed with multiple diseases, to select which disease you wish to add phenotype data for. If you do not see the “Add phenotype information to individual” option, then the individual does not have any diseases linked yet. In that case, edit the individual and select at least one disease from the list.

After you submitted the phenotype data, LOVD sends an email to the curator(s) and redirects you to the phenotype data overview.
10. Downloading & importing data

10.1 The LOVD3 import format

The LOVD3 import format is based on the LOVD2 tab-delimited import format, but all data is split into separate sections, preventing data duplication. Examples of sections are Genes, Transcripts, and Diseases. Each section has its own set of fields. Sections refer to each other using IDs. For instance, the Transcripts section has a geneid field that refers to an entry in the Genes section with the same value in its id field. This way, transcripts are connected to genes.

```plaintext
### LOVD-version 3000-010 ### Full data download ### To import, do not remove or alter this header ###
# charset = UTF-8
# max_errors = 50

## Genes ##

"{{id}}" "{{name}}" "{{chromosome}}" "{{chrom_band}}" "{{id_hgnc}}" "{{id_entrez}}" "{{id_omim}}"
"IVD" "isovaleryl-CoA dehydrogenase" "15" "q14-q15" "6186" "3712" "607036"

## Transcripts ##

"{{id}}" "{{geneid}}" "{{name}}" "{{id_ncbi}}" "{{id_protein_ncbi}}"
"00064" "IVD" "isovaleryl-CoA dehydrogenase, transcript variant 1" "NM_002225.3" "NP_002216.2"

## Diseases ##

"{{id}}" "{{symbol}}" "{{name}}" "{{id_omim}}"
"00006" "IVA" "isovaleric acidemia" "243500"

## Genes_To_Diseases ##

"{{geneid}}" "{{diseaseid}}"
"IVD" "00006"

## Individuals ##

"{{id}}" "{{owned_by}}" "{{statusid}}" "{{Individual/Lab_ID}}" "{{Individual/Reference}}"
"00000124" "00001" "9" "FB102" "{PMID:Vockley et al (2000):10677295}"

## Individuals_To_Diseases ##

"{{individualid}}" "{{diseaseid}}"
"00000124" "00006"

## Screenings ##

"{{id}}" "{{individualid}}" "{{owned_by}}" "{{Screening/Technique}}" "{{Screening/Template}}"
"000000116" "00000124" "00006" "RT-PCR;SEQ" "DNA;RNA"

## Screenings_To_Genes ##

"{{screeningid}}" "{{geneid}}"
"000000116" "IVD"
```

Figure 10.1: Example (partial) LOVD import format file. This example file contains information about the IVD gene, its first transcript variant, the IVA disease, the link between the IVD gene and the IVA disease, and an individual linked to the IVA disease and screened for variants in the IVD gene. Note the geneid field in the Transcripts section, which links a transcript to a gene, and the individualid field in the Screenings section, which links a screening to an individual. The link between genes and diseases and between screenings and genes are in a separate section because each gene can be linked to multiple diseases, and each screening can test for variants in multiple genes. Please note that a full LOVD download contains many more sections, and more columns. This file has been shortened to fit this page. This example file is therefore not a valid import file.

The first line of the file contains the file’s header line, which indicates the LOVD version number as well as the type of file. The LOVD version number is encoded and provided by
LOVD when creating a download. When importing, LOVD can decode the version and in case
the download format has changed between versions, convert the data into the correct format.
When this line is not included in the file that is imported, LOVD cannot process the file, and
halt.

Please NOTE that spreadsheets are well known for introducing errors in import-
ing/exporting text files, due to automatic interpretation of the values. When using
a spreadsheet program to edit a downloaded file, format all cells to "Text" before
importing/pasting the downloaded data.

Also some cases have been reported where MS Excel removed contents of large text
fields (> 255 characters).

The file’s data sections are indicated by lines starting with two hash characters (##),
followed by the name, again two hash characters, and a note that the header should not be
modified or removed. Other lines starting with two hash characters are regarded comments. A
downloaded file may contain comments, like the number of entries for each section.

The following sections are currently recognized:

Columns
Custom columns and their system-wide settings. In a full data download, this section
contains records created during installation of LOVD as well as records created by users.
During import, columns in the import file that already exist in the database are skipped.
In that case a warning is displayed, but the import will continue.

Genes
Genes, including external IDs. *Can currently not yet be imported.* When importing,
genomes can be listed in the file, but they should already be present in the database. If not
present, they should be created manually before importing the file. Links between genes
and diseases are described in the “Genes_To_Diseases” section.

Transcripts
Transcript variants, including external IDs. *Can currently not yet be imported.* When
importing, transcripts can be listed in the file, but they should already be present in the
database. If not present, they should be created manually before importing the file. A
transcript entry is linked to a gene using the “geneid” field.

Diseases
General disease information. Diseases should have a unique name and OMIM ID. During
import, diseases in the import file that already exist in the database (based on name or
OMIM ID) are skipped. In that case a warning is displayed, but the import will continue.
Links between diseases and genes are described in the “Genes_to_Diseases” section.

Genes_to_Diseases
Links genes and diseases by referring to their IDs.

Individuals *
Individuals and related information, like links to the parents of the individual. Links
between individuals and diseases are described in the “Individuals_to_Diseases” section.

Individuals_to_Diseases
Links individuals to diseases.
Phenotypes *
Phenotype information of a certain individual, related to a certain disease. A phenotype entry is linked to a disease using the “diseaseid” field. A phenotype entry is linked to an individual using the “individualid” field.

Screenings *
Variant screening information, with which variants are detected. A screening is linked to an individual using the “individualid” field. Links between screenings and genes are described in the “Screenings_to_Genes” section. Links between screenings and variants are described in the “Screenings_to_Variants” section.

Screenings_to_Genes
Linking screenings to genes, indicating which genes were screened using a certain screening.

Variants_on_Genome *
Genomic information of all variants detected. Links between variants and screenings are described in the “Screenings_to_Variants” section.

Variants_on_Transcripts *
Possible effects on a transcript caused by a genomic variant. A variant on transcript is linked to a variant on genome using the “id” field.

Screenings_to_Variants
Linking screenings to variants, indicating which variants were detected using a certain screening.

Categories indicated with an asterisk (*) are data types with custom columns (also see “Custom column categories”), and the exact list of fields in this category depend on the LOVD settings.

Lines in the import file starting with one hash character are settings. The following settings are currently recognized:

charset
Indicates which carset was used when creating the file. During import, usually the file encoding is detected given the file’s contents, in which case this setting is ignored.

max_errors
When this setting is not defined in the import file, LOVD stops processing an import file when 50 errors occurs. This default value can be modified in the import file with variable “max_errors”. Increasing the “max_errors” allows LOVD to process the import file without interruption. Decreasing the “max_errors” interrupts the import process earlier. Regardless of the value of “max_errors”, all errors must be solved before the import is saved to LOVD. Warnings however, do not interrupt the import process, and data can be saved in LOVD even when warnings are displayed.

Each section header is followed immediately by a line that indicates the names of the fields used in this section. When reading a downloaded file, this nicely indicates which column represents which data. When importing a file, being able to change this line allows you to skip certain columns or decide yourself in which order you would like to display the columns in the file. To clearly separate the line with column names from the data lines, the column names are surrounded by two curly braces on each side ({{ and }}). Field names can not contain spaces, match the field’s name in the database exactly, and the field names are separated by exactly one tab character from each other.
The section’s data is also in tab-delimited format. Each field is preferably enclosed by double quotes. All existing quotes in the data, are escaped by the backslash character (\), so " becomes \". Other special characters are \t for a tab character, and \r\n for a linebreak.

Specific fields explained

For each section, the import format contains all fields that are stored in the database. Some of these fields are not displayed on the normal LOVD data entry forms, and therefore may cause confusion. Others are displayed, but for the user appear as selection lists with human-readable values, whereas in the database only numbers are stored. To aid in understanding these fields and their values, all fields that may cause confusion are listed here.

Custom columns are not listed here, since every LOVD installation has different columns enabled and custom columns provide explanations about their use by themselves.

**allele** (from section Variants_On_Genome)
Indicates on which allele the variant is located. Does not necessarily imply inheritance!
This field cannot be updated during an import.

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Unknown</td>
<td>Heterozygous, parents not screened, or de novo and allele couldn’t be confirmed</td>
</tr>
<tr>
<td>1</td>
<td>Parent #1</td>
<td>Heterozygous, parents not screened, but useful in combination with ‘Parent #2’ to indicate cis or trans compound heterozygosity</td>
</tr>
<tr>
<td>2</td>
<td>Parent #2</td>
<td>Heterozygous, parents not screened, but useful in combination with ‘Parent #1’ to indicate cis or trans compound heterozygosity</td>
</tr>
<tr>
<td>3</td>
<td>Both (homozygous)</td>
<td>Variant is present on both alleles</td>
</tr>
<tr>
<td>10</td>
<td>Paternal (inferred)</td>
<td>Variant is on paternal allele, not confirmed in father but inferred from other information</td>
</tr>
<tr>
<td>11</td>
<td>Paternal (confirmed)</td>
<td>Variant is on paternal allele, confirmed in father</td>
</tr>
<tr>
<td>20</td>
<td>Maternal (inferred)</td>
<td>Variant is on maternal allele, not confirmed in mother but inferred from other information</td>
</tr>
<tr>
<td>21</td>
<td>Maternal (confirmed)</td>
<td>Variant is on maternal allele, confirmed in mother</td>
</tr>
</tbody>
</table>

If left empty during import, the default value is “0”.

**chromosome** (from sections Genes and Variants_On_Genome)
The chromosome related to this entry (gene or variant). When importing, do not fill ‘chr’ here as prefix, just the chromosome number or letter. This field cannot be updated during an import.

**chrom_band** (from section Genes)
The chromosome band, like ‘q15’.

**col_order** (from section Columns)
The order in which the columns of this column’s category are shown. Low values are shown before higher values.

**created_by** (from sections Columns, Genes, Transcripts, Diseases, Individuals, Phenotypes, Screenings and Variants_On_Genome)
The internal ID of the user that created this data entry. If left empty during import, the current user performing the import will be selected as the creator of this data entry.
Creators are allowed to edit the entry and add new data to the entry. They are fixed, and can not be re-assigned at a later time. This field cannot be updated during an import.

**created_date** (from sections Columns, Genes, Transcripts, Diseases, Individuals, Phenotypes, Screenings and Variants_On_Genome)
The date the entry was created, in the format YYYY-MM-DD HH:MM:SS. This field cannot be updated during an import.

**description_form** (from section Columns)
The column’s notes on the data entry form. This text is shown on the data entry form, just below the data field, for instance as clarification. See also section “Creating a new custom column”.

**description_legend_short** (from section Columns)
Description on short legend for this custom column. See also section “Creating a new custom column”.

**description_legend_full** (from section Columns)
Description on full legend for this custom column. See also section “Creating a new custom column”.

**diseaseid** (from sections Phenotypes)
ID referring to the ID in section Diseases. This field cannot be updated during an import.

**effectid** (from sections Variants_On_Genome and Variants_On_Transcripts)
The variant’s effect on a protein’s function. The first digit indicates the variant’s effect as reported by the source. The second digit indicates the variant’s effect as concluded by the curator. The possible values of each digit:

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-</td>
<td>Does not affect function, strong proof</td>
</tr>
<tr>
<td>3</td>
<td>-?</td>
<td>Probably does not affect function, strong evidence</td>
</tr>
<tr>
<td>5</td>
<td>?</td>
<td>Effect unknown</td>
</tr>
<tr>
<td>7</td>
<td>+?</td>
<td>Probably affects function, strong evidence</td>
</tr>
<tr>
<td>9</td>
<td>+</td>
<td>Affects function, strong proof</td>
</tr>
</tbody>
</table>

A score of 97 means a variant is reported to affect the function of the protein having strong proof, but the curator thinks more proof is needed, still accepting there is strong evidence for the variant’s effect. All options are:

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>ID</th>
<th>Name</th>
<th>ID</th>
<th>Name</th>
<th>ID</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>11</td>
<td>-/-</td>
<td>31</td>
<td>-?/-</td>
<td>51</td>
<td>?/-</td>
<td>71</td>
<td>+?/-</td>
</tr>
<tr>
<td>13</td>
<td>-/-?</td>
<td>33</td>
<td>-?/-?</td>
<td>53</td>
<td>?/-?</td>
<td>73</td>
<td>+?/-?</td>
</tr>
<tr>
<td>15</td>
<td>-/?</td>
<td>35</td>
<td>-/??</td>
<td>55</td>
<td>??/?</td>
<td>75</td>
<td>+?/?</td>
</tr>
<tr>
<td>17</td>
<td>-/+?</td>
<td>37</td>
<td>-?/+?</td>
<td>57</td>
<td>?/+?</td>
<td>77</td>
<td>+?/+?</td>
</tr>
<tr>
<td>19</td>
<td>-/+</td>
<td>39</td>
<td>-?/+</td>
<td>59</td>
<td>?/+</td>
<td>79</td>
<td>+?/+</td>
</tr>
</tbody>
</table>

In the Variants_On_Transcripts section, this field indicates the effect of the variant on this specific transcript. In the Variants_On_Genome section, this field indicates the overall effect of the variant on the function of any gene it influences; basically, the maximum score of all of the variant’s effectid values. If left empty during import, the default value will be “55”.

**fatherid** (from section Individuals)
Optional; The internal ID of the individual in the database that is the father of this individual.
The LOVD3 import format

form_type (from section Columns)
The custom column’s form type definition. When importing new custom columns, we recommend you use the Data type wizard available when creating a new custom column, and to use the form type definition generated by the wizard. See also section “Creating a new custom column”.

head_column (from section Columns)
The custom column’s header, shown above the column in data views. See also section “Creating a new custom column”.

hgvs (from section Columns)
If set to 1, this column is HGVS recommended. These columns can not be removed. This setting can not be set when importing columns. Default value is 0.

id_entrez (from section Genes)
This field cannot be updated during an import.

id_hgnc (from section Genes)
This field cannot be updated during an import.

id_mutalyzer (from section Transcripts)
This field cannot be updated during an import.

id_ncbi (from section Transcripts)
This field cannot be updated during an import.

id_omim (from section Genes)
This field cannot be updated during an import.

id_protein_ncbi (from section Transcripts)
This field cannot be updated during an import.

imprinting (from section Genes)
Whether or not a gene is imprinted (either maternal or paternal).

individualid (from sections Screenings and Phenotypes)
ID referring to the ID in section Individuals. This field cannot be updated during an import.

mandatory (from section Columns)
If set to 1, this column is mandatory. When entering new data, this custom column’s field must be filled in. See also section “Creating a new custom column”. Default value is 0.

mapping_flags (from section Variants_On_Genome)
Information about how LOVD handles the mapping of this variant. LOVD has an automatic mapping script that finds transcripts in the range of a genomic variant, and can annotate the genomic variant on the transcript using the Mutalyzer web service. Whether or not LOVD is allowed to map the variant, and how LOVD will handle the mapping, is stored in this field. This field is a boolean flag column; multiple options can be grouped in this field by adding the numbers up.
### Possible values:

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No mapping allowed</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>MAPPING_ALLOW</td>
<td>When set, the variant can be mapped by the LOVD mapping script</td>
</tr>
<tr>
<td>2</td>
<td>MAPPING_ALLOW_CREATE_GENES</td>
<td>When set, LOVD is allowed to create new genes in the database if needed to annotate the variant</td>
</tr>
<tr>
<td>4</td>
<td>MAPPING_IN_PROGRESS</td>
<td><em>Not used when importing</em>; when set, LOVD is currently busy mapping the variant</td>
</tr>
<tr>
<td>8</td>
<td>MAPPING_NOT_RECOGNIZED</td>
<td><em>Not used when importing</em>; when set, LOVD did not recognize the variant, and was therefore unable to map the variant</td>
</tr>
<tr>
<td>16</td>
<td>MAPPING_ERROR</td>
<td><em>Not used when importing</em>; when set, LOVD has encountered an error while mapping</td>
</tr>
<tr>
<td>32</td>
<td>MAPPING_DONE</td>
<td><em>Not used when importing</em>; when set, LOVD is done mapping the variant, regardless of whether the variant actually got mapped</td>
</tr>
</tbody>
</table>

For instance, a value of 11 means for this variant mapping has been enabled (1), LOVD is allowed to create new genes while mapping the variant (2), but LOVD does not recognize the variant and therefore can not map it (8). $1 + 2 + 8 = 11$. If left empty during import, the default value will be “0”.

**motherid** (from section Individuals)
Optional; The internal ID of the individual in the database that is the mother of this individual.

**mysql_type** (from section Columns)
The custom column’s MySQL data type. When importing new custom columns, we recommend you use the Data type wizard available when creating a new custom column, and to use the MySQL data type generated by the wizard. See also section “Creating a new custom column”.

**note_index** (from section Genes)
Notes that should appear on the gene home page.

**note_listing** (from section Genes)
Notes that should appear on gene-specific data listings.

**owned_by** (from section Individuals, Phenotypes, Screenings and Variants_On_Genome)
Optional; The internal ID of the user that owns this data entry. Owners are allowed to edit the entry and add new data to the entry. Owners are not fixed, and can be re-assigned. When no value is provided during import, this value is set to the user ID of the user importing the data.

**panelid** (from section Individuals)
Optional; The internal ID of the individual in the database that represents the group to which this individual or group of individuals belong to.
panel_size (from section Individuals)
The number of individuals represented by this entry (default: 1). Only used when multiple individuals, for instance from one family, are aggregated into one entry in the database.

position_c_cds_end (from section Transcripts)
The position, relative to the translation start site, of where the translation ends. Is always a positive whole number, should always be dividable by three. This field cannot be updated during an import.

position_c_mrna_start (from section Transcripts)
The position, relative to the translation start site, of where the transcript starts. Is always a negative whole number. This field cannot be updated during an import.

position_c_mrna_end (from section Transcripts)
The position, relative to the translation start site, of where the transcript ends. Is always a positive whole number. This field cannot be updated during an import.

position_c_start (from section Variants_On_Transcripts)
The position, relative to the translation start site, of the first base of the variant. The first base of the variant is defined by the HGVS nomenclature rules, and may be the same as the last base. This field cannot be updated during an import.

position_c_start_intron (from section Variants_On_Transcripts)
The position, relative to the start or end (whichever is closest) of the intron, of the first base of the variant. The first base of the variant is defined by the HGVS nomenclature rules, and may be the same as the last base. This field only applies to variants that start in an intron. This field cannot be updated during an import.

position_c_end (from section Variants_On_Transcripts)
The position, relative to the translation start site, of the last base of the variant. The last base of the variant is defined by the HGVS nomenclature rules, and may be the same as the first base. This field cannot be updated during an import.

position_c_end_intron (from section Variants_On_Transcripts)
The position, relative to the start or end (whichever is closest) of the intron, of the last base of the variant. The last base of the variant is defined by the HGVS nomenclature rules, and may be the same as the first base. This field only applies to variants that end in an intron. This field cannot be updated during an import.

position_g_mrna_start (from section Transcripts)
The genomic position, relative to the chromosome, of where the transcript starts. This field cannot be updated during an import.

position_g_mrna_end (from section Transcripts)
The genomic position, relative to the chromosome, of where the transcript ends. This field cannot be updated during an import.

position_g_start (from section Variants_On_Genome)
The genomic position, relative to the chromosome, of the first base of the variant. The first base of the variant is defined by the HGVS nomenclature rules, and may be the same as the last base. This field cannot be updated during an import.

position_g_end (from section Variants_On_Genome)
The genomic position, relative to the chromosome, of the last base of the variant. The...
last base of the variant is defined by the HGVS nomenclature rules, and may be the same
as the first base. This field cannot be updated during an import.

**preg_pattern** (from section Columns)
For text field custom columns, regular expression checks can be performed. See also
section “Creating a new custom column”.

**public_view** (from section Columns)
If set to 1, this column’s contents are visible to the public (only for public entries). See
also section “Creating a new custom column”.

**public_add** (from section Columns)
If set to 1, submitters can fill in this field. Otherwise, only curators and higher level users
can enter data in this field.

**refseq_genomic** (from section Genes)
The genomic reference sequence, this field cannot be updated during an import.

**refseq_UD** (from section Genes)
This is an internal ID for the genomic reference sequence, generated by Mutalyzer. This
field cannot be updated during an import.

**select_options** (from section Columns)
When the custom column is a selection field, the selection options are listed here. See also
section “Creating a new custom column” for the format. Please remember that linebreaks
in the LOVD import format are displayed as \r\n.

**standard** (from section Columns)
If set to 1, this column is enabled by default. When importing, this is only useful for
shared custom columns. When set, the column will then be enabled by default for a new
disease or gene, depending on the column category. See also section “Creating a new
custom column”. Default value is 0.

**statusid** (from section Individuals, Phenotypes and Variants_On_Genome)
Optional; This numeric status ID defines if the data is public or not.

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>In progress</td>
<td><em>Not used when importing;</em> data entry is in the process of being submitted</td>
</tr>
<tr>
<td>2</td>
<td>Pending</td>
<td>Data entry has been submitted, awaiting approval of curator</td>
</tr>
<tr>
<td>4</td>
<td>Non public</td>
<td>Data entry is not public, hidden from view for non-authorized users</td>
</tr>
<tr>
<td>7</td>
<td>Marked</td>
<td>Data entry is public but marked, shows up as red for authorized users</td>
</tr>
<tr>
<td>9</td>
<td>Public</td>
<td>Data entry is public</td>
</tr>
</tbody>
</table>

Data with status “In progress”, value 1, can not be imported. Instead, there are 3 options:

1. Ask the submitter to finish the submission.
2. Change the statusid value to a valid value, being 4 or higher.
3. Exclude this record for import, by either removing it from the import file, or hiding
   the record by putting a # in the beginning of the line. If you do this, make sure
   that all related data from other sections are excluded as well, or LOVD will return
   errors when trying to resolve references in the file.
When no value is provided during import, this value is set to ‘9’ (Public).

**symbol** (from section Diseases)
The disease’s abbreviation, preferably the most commonly used one.

**transcriptid** (from section Variants_On_Transcripts)
ID referring to the ID in section Transcripts. This field cannot be updated during an import.

**type** (from section Variants_On_Genome)
The type of variant on DNA level, as defined by the Mutalyzer web service. All possible values are:

<table>
<thead>
<tr>
<th>ID</th>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>comp</td>
<td>Compound variant</td>
<td>Not normally in use; multiple variants grouped together</td>
</tr>
<tr>
<td>del</td>
<td>Deletion</td>
<td>One or more DNA bases have been deleted</td>
</tr>
<tr>
<td>delins</td>
<td>Deletion/Insertion</td>
<td>One or more DNA bases have been deleted, while others have been inserted; when not a substitution</td>
</tr>
<tr>
<td>dup</td>
<td>Duplication</td>
<td>Duplication of one or more DNA bases</td>
</tr>
<tr>
<td>ins</td>
<td>Insertion</td>
<td>Insertion of one or more DNA bases; when not a duplication</td>
</tr>
<tr>
<td>subst</td>
<td>Substitution</td>
<td>One DNA base has been replaced by another</td>
</tr>
</tbody>
</table>

**updated_by** (from section Genes)
The internal ID of the user who was the last to modify data that is linked to the gene. This field cannot be updated during an import.

**updated_data** (from section Genes)
The date data linked to the gene was modified last. This field cannot be updated during an import.

**variants_found** (from section Screenings)
If set to 1, this screening is reported to have found variants. If set to 0, this screening is reported to not have found variants. Please note that other values are not allowed.

**width** (from section Columns)
The displayed column width in pixels. See also section “Creating a new custom column”.

### 10.2 Downloading & importing data

#### Downloading data from LOVD

For backup purposes or to move data sets from one LOVD into another, LOVD supports different download options. All downloaded file types lack the LOVD system settings, registered user accounts, and object-specific custom column settings.

In order to have a full backup, for instance to restore LOVD in case of a hardware failure, more information is needed. It is therefore recommended not to rely only on this feature for a full backup of LOVD. However, it works perfectly as a backup for your variant and patient data, for instance if you accidentally deleted the wrong entry or the wrong column.

If you’re planning large parallel changes to data entries in LOVD using spreadsheet software such as OpenOffice.org Calc or MS Excel, you could use the download files as a starting point.
In that case, read the instructions at “Downloading & importing the complete data” before you begin.

**Importing data sets into LOVD**

For managers it is possible to import data directly into LOVD, using the LOVD3 import format. For curators, only simulation is available (see below), starting from LOVD 3.0-14. See section “The LOVD3 import format” for more details about the file format. Files created or modified using spreadsheet software such as OpenOffice.org Calc or MS Excel must be saved as a text file. The file must be tab delimited, otherwise LOVD will produce an error.

To import a dataset to LOVD, click the “Import data” link from the “Setup” tab dropdown menu, or from the setup area, click the “Import data using the LOVD import format” link listed under “Download & Import”. Curators can import data from the Configuration area. Click the “Import data using the LOVD import format” link listed under “Download & Import”.

The LOVD import has a simulate option, allowing you to check if the import file is correct, without importing anything. Check the “Simulate (don’t actually import the data)” checkbox to use this simulation option. *This option is always enabled for curators, and cannot be disabled.* This allows curators to test their import files, but it still requires a manager or up to actually import the file.

On the import form, in the “Import options” section, you’ll find the “Import mode”. The two options available are “Update existing data” (see section “Update existing data via import”) and “Add only, treat all data as new” (see section “Insert new data via import”).

**Insert new data via import**

The option “add only” on the import page will insert all data in the import file as new, also if the identifiers used are already present in the database. See section “The LOVD3 import format” for more details about the file format.

**Update existing data via import**

For managers and higher it is possible to update certain data via an import (curators can run a simulate). There are some restrictions on what you can modify and what not. In general, when you can modify certain data via the LOVD data entry forms, you can modify them via the update import as well. Furthermore, to prevent unintended edits, you can currently only modify one field per line.

You cannot modify the created_by, created_date, edited_by, edited_date fields (from sections Columns, Genes, Transcripts, Diseases, Individuals, Phenotypes, Screenings and Variants_On_Genome). These fields are set by LOVD and changes in these fields are ignored. See the section “Specific fields explained” to check which fields you are not allowed to update.

The IDs in the import file are used to identify the records in the LOVD database. Therefore, you cannot modify the ID fields in your import file. To make sure your IDs in the LOVD database correspond with the IDs in your import file, and to prevent unintended edits, it is recommended to make a new download (complete data set, custom columns, own data set or a gene’s data set, depending on what you would like to update) before you start preparing your changes.
On the import form, select the “Update existing data” option for the “Import mode” field, if you wish to modify data in your LOVD database. You can use the simulate option to check your file for errors first.

10.3 Downloading & importing the complete data

Downloading the complete data set from LOVD

LOVD supports a full download that contains most (currently not all) data stored in LOVD. The data sections included in the download file are: Columns, Genes, Transcripts, Diseases, Genes_To_Diseases, Individuals, Individuals_To_Diseases, Phenotypes, Screenings, Screenings_To_Genes, Variants_On_Genome, Variants_On_Transcripts and Screenings_To_Variants. In order to have a full backup, for instance to restore LOVD in case of a hardware failure, more information is needed. The download file lacks the LOVD system settings, registered user accounts, and object-specific custom column settings. It is therefore recommended not to rely only on this feature for a full backup of LOVD. However, it works perfectly as a backup for your variant and patient data, for instance if you accidentally deleted the wrong entry or the wrong column.

To download the full dataset from LOVD, click the “Download all data” link from the “Setup” tab dropdown menu, or from the setup area, click the “Download all data in LOVD import format” link listed under “Download & Import”. The download begins immediately. Depending on your browser, it will save the file to a default location on your hard disk, or ask you whether to open or to save the file.

Importing the complete data set into LOVD

A complete dataset downloaded from the LOVD can be imported into a LOVD. This can only be done by the manager. Not all data can and will be imported and some data needs to be modified to be able to import in LOVD. Genes and transcripts can currently not be imported directly. See section “The LOVD3 import format” for details on each section.

10.4 Downloading & importing the custom column data

Downloading custom columns data from LOVD

Managers can download a data file with only the custom columns which are added to an LOVD installation. This can be done for backup purposes, but also for importing to another LOVD installation. The records downloaded can be both records created during installation of LOVD, as well as records created by users. To download the custom columns data, click the “Download all LOVD custom columns” link from the setup menu tab dropdown menu, or from the setup area, click the “Download all LOVD custom columns in the LOVD import format” link listed under “Custom data columns”. The download begins immediately. Depending on your browser, it will save the file to a default location on your hard disk, or ask you whether to open or to save the file.
Importing a custom columns data set into LOVD

The custom column data set downloaded by a manager can be uploaded to the same or any other LOVD. Columns which already exists in the database are not imported, but skipped instead. For this, LOVD checks if the column’s ID already exists in the database. No other fields are compared.

10.5 Downloading & importing own data

Downloading own data set from LOVD

LOVD allows you to collect a file containing all information in LOVD that has been submitted by you, or is currently owned by you. Proceed to your account details by clicking the “Your account” link on the top right-hand side of the screen. Then, click the “Options” button and click “Download all my data”. The download begins immediately. Depending on your browser, it will save the file to a default location on your hard disk, or ask you whether to open or to save the file.

This file, compared to the full data download that LOVD offers to managers and the database administrator, lacks the Columns data section. The following sections contain data that is not owned by the user in question, but merely serve as a reference to allow interpretation of all data: Genes and Transcripts (the ones where variants were reported in are shown), and Diseases (the ones linked to submitted Individuals are shown). These section are marked with “## For reference only, not part of the selected data set”.

Importing own data set into LOVD

The own data set downloaded by a submitter can also be reuploaded into LOVD. But as with all import files, only a manager can import the data file.

If the diseases in the import file do not exist in LOVD, these diseases will be added during the import. Downloads in this format do not contain any links between genes and diseases. To create this link, you will have to add records to the “Genes_To_Diseases” section, or manually create the links. See “Relation to diseases” for how to create a link between a gene and a disease.

For more details on each section of the import file, see section “The LOVD3 import format”.

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10.6 Downloading & importing a gene’s data

Downloading a gene’s data set from LOVD

Curators can download a file containing all information related to a gene they have authorization on. This can be done for backup purposes, but also for importing to an other LOVD installation. From the gene homepage, click the “Options” button and click “Download all this gene’s data”. The download begins immediately. Depending on your browser, it will save the file to a default location on your hard disk, or ask you whether to open or to save the file.

This file, compared to the full data download that LOVD offers to managers and the database administrator, lacks the Columns data section. All sections are filtered to show only data related to the gene in question. The following sections are perhaps not obvious: Diseases (the ones related to the gene in question, as well as the ones linked to the individuals in this file, are shown), Phenotypes (the ones linked to the individuals in this file are shown), and Screenings (the ones which found variants in the gene is question are shown). The individuals in this file may have more variants reported, which are not affecting this gene and therefore not shown in this file.

Importing a gene’s data set into LOVD

The gene data set downloaded can be uploaded without modifications into the same or any other LOVD installation. However, the gene and its transcript(s) must exist in the database in which the file is imported. The column section is missing in this download file, but apart from that, this is the same import file as the complete data set. See section “Importing the complete data set into LOVD” on details about importing a gene’s data set.
11. The LOVD APIs

LOVD3 contains an LOVD2-style data retrieval API, useful for simple queries and retrieval of basic gene and variant information, and a new data submission API, introduced in LOVD 3.0-21, useful for submitting data from other software tools without the need of using the web forms.

11.1 Data retrieval API

Just like LOVD2 (from version 2.0-22, released October 5th 2009), LOVD3 includes a simple webservice enabling simple queries or listing of variant data (not patient data). This API only shows very basic information and can therefore not be disabled. The LOVD installations that have indicated to be included in the global LOVD listing are included in the overall LOVD querying service, which is based on this API.

Examples of possibilities are searching on a gene symbol, getting the list of available genes in the database, or on a per-gene basis, list all variants or search for a certain variant or DNA location. Even though LOVD3 is not gene-based like LOVD2 was, we have decided to keep the same gene-centered design of the API to make sure LOVD2 and LOVD3 can be queried alike. The output it creates is either an Atom 1.0 feed with the LOVD information in plain text, or a JSON file.

A more advanced API will be included in LOVD3 later, allowing the querying of variants genome-wide and getting more detailed information out of LOVD, such as patient and phenotype data. This API will be configurable, so submitters decide for themselves under which data sharing license which type of information will be shared.

Terms of service and fair use policy

These terms of service and fair use policy are applicable to all APIs present on the LOVD site and any LOVD installations hosted by the Leiden University Medical Center, recognizable by domain names ending by .lovd.nl or .lumc.nl. Other servers hosting LOVDs may also have terms of service in place. If in doubt, try to contact the party hosting the LOVD installation.

- Read the API possibilities below carefully, and choose the most efficient method to query the API — meaning, the least amount of requests to get the information you need.
- Limit your requests to a maximum of 5 per second per server/domain name.
- If the API indicates a certain entry does not exist, do not repeat the same request for at least 4 hours, unless you have reason to believe the entry may exist now.
- Not required, but appreciated; please set a ‘User agent’ string that identifies you, so we can see how the service is being used and we can contact you if necessary.
The LOVD APIs

Data retrieval API

Format

Since 3.0-22, the LOVD2-style API allows for JSON data to be returned. The original format, also still available in 3.0-22, is an Atom feed, and is still the standard. When searching for genes, an Atom feed is returned with zero or more entries. When requesting a specific entry, just the Atom entry is returned. The Atom format is described in detail on Wikipedia.

The content of the Atom entries is in plain text, with headers and values separated by a colon (:). Other formats as a ‘payload’ of the Atom entries are not available.

The new LOVD3 API (release date unavailable) will probably support multiple formats, such as VarioML and JSON, and will not support Atom feeds, since this API serves a different purpose.

Genes

```
<?xml version="1.0" encoding="UTF-8"?>
<feed xmlns="http://www.w3.org/2005/Atom">
  <title>
    Results for your query of the database
  </title>
  <link rel="alternate" type="text/html" href="http://databases.lovd.nl/shared/>
  <link rel="self" type="application/atom+xml" href="http://databases.lovd.nl/shared/api/rest/genes/>
  <updated>2018-04-17T20:15:10+02:00</updated>
  <id>tag:databases.lovd.nl,2012-05-02:web01:shared/REST_api</id>
  <generator uri="http://www.LOVD.nl/" version="3.0-21">
    Leiden Open Variation Database
  </generator>
  <rights>Copyright (c), the curators of this database</rights>
  <entry xmlns="http://www.w3.org/2005/Atom">
    <title>IVD</title>
    <link rel="alternate" type="text/html" href="http://databases.lovd.nl/shared/genes/IVD/>
    <link rel="self" type="application/atom+xml" href="http://d...lovd.nl/shared/api/rest/genes/IVD/>
    <id>tag:databases.lovd.nl,2011-04-05:IVD</id>
    <author>
      <name>Gerard C.P. Schaafsma</name>
    </author>
    <contributor>
      <name>Ivo F.A.C. Fokkema</name>
    </contributor>
    <published>2011-04-05T16:04:49+02:00</published>
    <updated>2017-08-07T13:40:16+02:00</updated>
    <content type="text">
      id:IVD
      entrez_id:3712
      symbol:IVD
      name:isovaleryl-CoA dehydrogenase
      chromosome_location:15q14-q15
      position_start:chr15:40697686
      position_end:chr15:40713512
      refseq_genomic:NG_011986.1
      refseq_mrna:NM_002225.3
      refseq_build:hg19
    </content>
  </entry>
</feed>
```

Figure 11.1: Example LOVD API output, in the default Atom format. The longest line has been shortened to fit this page. This example output of a search on gene symbol returns the IVD gene and its basic information. Source: LOVD3 shared installation.

The gene information shared through the API consists of the official gene symbol (**id**, **symbol**), the gene’s Entrez ID (**entrez_id**), the gene’s official name (**name**), the chromosome...
11. The LOVD APIs

Data retrieval API

Figure 11.2: Example LOVD API output, in the JSON format, of the same API search result as the previous figure. Note the slight differences in the data returned; the JSON format adds the chromosome field, returns all transcripts associated with the gene in question, and stores the curator information in an array. Source: LOVD3 shared installation.

band (chromosome_location) and exact locations (position_start, position_end, not always available), its genomic reference sequence (refseq_genomic), first transcript added (refseq_mrna, all transcripts when using JSON) and the code of the reference sequence build that the genomic positions are based on (refseq_build). The JSON data has slightly more fields (created_by, created_date, curators, updated_date), which are also available in the Atom entry itself (not the plain text payload).

The official gene symbol is mentioned twice, because in LOVD2 multiple databases could exist for the same gene by setting a suffix on the gene symbol; there the ‘id’ field could have a different value than the ‘symbol’ field. Having two fields for the gene symbol is retained in the API to keep the format between LOVD3 and LOVD2 consistent.

Please note, also for consistency with the LOVD2 API, that in the Atom entry, the ‘refseq_mrna’ field can contain only one value whilst in LOVD3 one gene can contain multiple transcripts. The Atom entry shows only the first transcript, sorted on their NCBI ID. The JSON data shows all transcripts.
11. The LOVD APIs

Data retrieval API

Variants

<?xml version="1.0" encoding="UTF-8"?>
<feed xmlns="http://www.w3.org/2005/Atom">
  <title>
    Results for your query of the IVD gene database
  </title>
  <link rel="alternate" type="text/html" href="http://databases.lovd.nl/shared/">
  </link>
  <link rel="self" type="application/atom+xml" href="http://...lovd.nl/shared/api/rest/variants/IVD">
  </link>
  <updated>2017-08-07T13:40:16+02:00</updated>
  <id>tag:databases.lovd.nl,2012-05-02:web01:shared/REST_api</id>
  <generator uri="http://www.LOVD.nl/" version="3.0-21">
    Leiden Open Variation Database
  </generator>
  <rights>Copyright (c), the curators of this database</rights>
  <entry xmlns="http://www.w3.org/2005/Atom">
    <title>IVD:c.860G>A</title>
    <link rel="alternate" type="text/html" href="http://...?search_VariantOnGenome/DBID=IVD_000001"/>
    <link rel="self" type="application/atom+xml" href="http://.../api/rest/variants/IVD/0000000563"/>
    <id>tag:databases.lovd.nl,2011-04-06:IVD/0000000563</id>
    <author>
      <name>Gerard C.P. Schaafsma</name>
    </author>
    <contributor>
      <name>Ivo F.A.C. Fokkema</name>
    </contributor>
    <published>2011-04-06T16:20:25+02:00</published>
    <updated>2017-03-28T16:27:49+02:00</updated>
    <content type="text">
      symbol:IVD
      id:0000000563
      position_mRNA:NM_002225.3:c.860
      position_genomic:chr15:40707154
      Variant/DNA:c.860G>A
      Variant/DBID:IVD_000001
      Times_reported:1
    </content>
  </entry>
</feed>

Figure 11.3: Example LOVD API output, in the default Atom format. The three longest lines have been shortened to fit this page. This example output of a search on DB ID returns one variant in the IVD gene and its basic information. Source: LOVD3 shared installation.

The variant information shared through the API consists of the official gene symbol of the selected gene (symbol), the variant’s internal ID (id), the variant’s position relative to the transcript in HGVS-like format (position_mRNA), the variant’s genomic position in HGVS-like format (position_genomic), the DNA field in HGVS format, HTML encoded in the Atom format to not break XML parsers (Variant/DNA), the variant’s DBID value (Variant/DBID) and the panel size of the individual of the variant entry (Times_reported). The JSON data has slightly more fields (owned_by, created_by, created_date, edited_date), which are also available in the Atom entry itself (not the plain text payload). Also, the JSON data will list the variant’s position and the variant’s DNA change on transcript level for all transcripts that the variant is mapped on.

If the variant could not be interpreted, the position fields will contain a question mark. The difference between the ‘id’ field and the ‘Variant/DBID’ field is that the first is the ID of the variant observation, whilst the latter is the ID unique for this variant, but shared for each observation. It is used link back to LOVD from genome browsers, for instance. When
variant observations are grouped together, the ‘Times_reported’ field will contain the sum of the values of all observations together.
Possibilities

A full list of options for this API is included below. Please note that although “rest.php” is included in all the URLs, simply “rest” will work also. However, for LOVD2 installations, this does not always work, and as such the extension is included. To obtain JSON output, add the ‘format=application/json’ flag to all URLs.

Genes

Listing of all genes in the database
http://databases.lovd.nl/shared/api/rest.php/genes

Searching on the gene symbol (full match only)
http://databases.lovd.nl/shared/api/rest.php/genes?search_symbol=IVD

Showing only one specific gene entry
http://databases.lovd.nl/shared/api/rest.php/genes/IVD

Searching on the genomic position

Chromosome only:
http://databases.lovd.nl/shared/api/rest.php/genes?search_position=chr15

Chromosomal location:
http://databases.lovd.nl/shared/api/rest.php/genes?search_position=chr15:40705600

Chromosomal range, exact match (only match genes having exactly this range)
http://databases.lovd.nl/shared/api/rest.php/genes?
    search_position=chr15:40697686_40713512&position_match=exact

Chromosomal range, exclusive match (only match genes completely within this range)
http://databases.lovd.nl/shared/api/rest.php/genes?
    search_position=chr15:40657686_40762512&position_match=exclusive

Chromosomal range, partial match (match any gene overlapping the given region)
http://databases.lovd.nl/shared/api/rest.php/genes?
    search_position=chr15:40657686_40762512&position_match=partial
Variants

Listing of all variant entries in a certain gene
http://databases.lovd.nl/shared/api/rest.php/variants/IVD

Searching on the DNA position

Coding DNA or genomic position, exact match only
This does not allow for partial matches, so variant c.157_158del is not matched. c.157 will match c.157+? and c.157_158 will match c.157+?_158-?. However, c.157 does not match c.157+5. Searching on genomic locations can be achieved using g. as a prefix.
http://databases.lovd.nl/shared/api/rest.php/variants/IVD?search_position=g.40699840

Genomic position only, exclusive match (only match variants completely within this range)
http://databases.lovd.nl/shared/api/rest.php/variants/IVD?
  search_position=g.40710300_40710410&position_match=exclusive

Genomic position only, partial match (match any variant overlapping the given region)
http://databases.lovd.nl/shared/api/rest.php/variants/IVD?
  search_position=g.40710300_40710410&position_match=partial

Searching on the DNA field
http://databases.lovd.nl/shared/api/rest.php/variants/IVD?
  search_Variant/DNA=c.157C>T
This does not allow for partial matches, but c.(157C>T) or c.157C>T? will also match.

Searching on the DBID field
http://databases.lovd.nl/shared/api/rest.php/variants/IVD?
  search_Variant/DBID=IVD_0000001

Showing only one specific variant entry (internal ID only)
http://databases.lovd.nl/shared/api/rest.php/variants/IVD/0000001350

World-wide LOVD querying service

There is also a world-wide LOVD query API that allows you to query a genomic location to find out whether there is a variant reported in any public LOVD installation that has registered at LOVD.nl. Please contact us directly for more information on that API.
11.2 Data submission API

LOVD3 includes a data submission API, useful for submitting JSON data files directly into LOVD3 without the need of web forms. This is especially useful for diagnostic labs that wish to submit their found variants to LOVD3 without the hassle of having to submit them manually or to construct LOVD3 import files.

Note that this API was introduced in LOVD 3.0-18, but was only fully functional from LOVD 3.0-21.

The submission format

The format used by the submission API is a JSON implementation of the VarioML format. For an example file, please see http://LOVD.nl/3.0/submission_api_format.json. Since VarioML is originally an XML format, all XML attributes have been prefixed by an '@' character, and text nodes are stored in '#text' elements. The JSON format allows for multiple individuals to be submitted at once; the file is required to contain at least one individual with at least one variant. The mandatory columns are the same as otherwise mandatory in LOVD3. For fields that require a fixed set of options, such as the individual’s gender, the possible options are mentioned in a comment in the example JSON file. JSON officially can’t contain comments so a JSON parser might fail on the example file, but LOVD3 is OK with them.

Do note that not all fields that may be configured in LOVD3 will be representable (yet) in this format. All fields currently supported are present in the example file (see above). If you have more data that you’d like to submit using this format, please contact us. We aim to map as many LOVD3 fields as possible to VarioML.

Mandatory fields to set before submission

Before you can submit your data using this example file, you’ll have to change the values of some of the fields. The ‘lsdb’ element has an ‘@id’ and an ‘@uri’ field that needs to be set. The ‘@id’ is the LOVD’s ID, which can be found by the database administrator in the API authorization token dialog (see “Getting your authorization token”. If you do not know who the LOVD’s administrator is, submit the file without a valid ID (see “Submitting the JSON file”) and LOVD will respond with the administrator’s contact information.

The ‘source’ element under ‘lsdb’ also requires a few fields to be filled in. For ‘name’, fill in the name of the lab that is submitting the data. Under ‘contact’, fill in your name under ‘name’, and your email under ‘email’. The ‘db_xref’ fields are also required, and are explained below.

Getting your authorization token

To submit data to LOVD3 using the submission API, you need to create an authorization token for your submitter account. To do so, log into LOVD3 and proceed to your account page by clicking the “Your account” link in the top right side of the screen. In the “API token” field, click “Show / More information”. In the new dialog, click the “Create new token” button to create a token. Select the desired validity and click “Create new token”. If the token was created successfully, click “Back” and copy your token from the bottom of the information dialog (see figure 11.5). In the JSON data, under ‘lsdb’ / ‘source’ / ‘contact’ / ‘db_xref’, fill in both ‘@accession’ fields. Where ‘@source’ is ‘lovd’, fill in your numeric account ID.
(on the top of your account page). Where ‘@source’ is ‘lovd_auth_token’, fill in your created token.

![API authorization token](image)

**Figure 11.5:** The authorization token dialog, showing your current token at the bottom and including buttons to create a new token or just revoking the current one. The database administrator also can find the LOVD’s ID at the top of this dialog.

### Submitting the JSON file

To submit your JSON file, **POST** it to `/api/submissions` for the intended LOVD instance. In case the intended LOVD is located at [https://databases.lovd.nl/shared/](https://databases.lovd.nl/shared/), then **POST** your file to [https://databases.lovd.nl/shared/api/submissions](https://databases.lovd.nl/shared/api/submissions). Before submitting though, please read the information under “Different versions of this API”. See figure 11.6 for an example on how to submit using `curl`.

```bash
curl --data @submission.json "https://databases.lovd.nl/shared/api/submissions"
```

**Figure 11.6:** Example of how to submit a VarioML JSON file to the submission API, using `curl`. In this example, the JSON file is named `submission.json` and is located in the current working directory.

The submission API will respond in JSON, sending its version, a messages array, a warnings array, an errors array, and a data array (see figure 11.7).

The ‘messages’ array is for information of successful submissions. The ‘warnings’ array is for output of information you should review because it might be a potential problem, although it will not prevent a successful submission. The ‘errors’ array contain fatal errors. The ‘data’ array is reserved for returning data and not used in **POST** requests.

A successful submission results in an LOVD3 import file being created on the server. This file then needs to be imported into LOVD3. In LOVD 3.0-21, submissions done through the API are not yet scheduled for import automatically, it requires manual intervention of a manager. See the INSTALL.txt file that is shipped with LOVD for more information.
11. The LOVD APIs

Data submission API

```json
{
    "version": 1,
    "messages": [],
    "warnings": [],
    "errors": [
        "VarioML error: LSDB root element not found, or has no ID."
    ],
    "data": []
}
```

**Figure 11.7:** Example submission API output, when sending `{"lsdb":{}}`. All arrays shown are always present, but can be empty.

Different versions of this API

Before you submit, make sure you read this information, to prevent unexpected problems if we would update the API.

To allow easy further development of this API, we might change the way the API works or change the way the API returns data. To make sure this doesn’t harm your application, you can instruct LOVD to use a fixed version of the API. To do so, you must include the APIs version in the URL. For instance, to always use version 1 of the API, even when version 2 is already released, use `/api/v1/submissions` instead of `/api/submissions`. When you’re not supplying a version number in the URL you POST to, LOVD will automatically use the latest version of the API. Make a decision based on what works best for you in your situation.
12. Keeping your data secure

LOVD contains several important security measures to protect your data from unauthorized access. This page explains IT experts what kind of security measures are implemented in LOVD, but it also shows what you as a LOVD user can do to protect your data.

12.1 Built-in security measures

After a public request on a programmer’s internet forum a LOVD installation had been subjected to various attacks and hack attempts. Furthermore, the source code has been security audited by three different software packages to ensure no security holes were left open. Some of the security risks LOVD protects you against, are displayed below.

Safe login procedure

If someone tries to break into LOVD by repeatedly trying different passwords (brute-forcing) for a correct username, this account will be locked, preventing unauthorized access. The account can be unlocked by a manager or the database administrator. They can also check the LOVD logs which IP address was responsible of the locking of the account, which can be used by the server’s system administrator to block access to LOVD from this IP address entirely.

As an additional security measure, each user can restrict access to their account from only a certain IP address or a range of IP addresses, preventing unauthorized access, even with the correct password, from unauthorized IP addresses.

Protection against SQL injection

SQL injection is the process of including unwanted SQL database commands into a database through the manipulation of online web forms. SQL injection can be used for, amongst other things, unauthorized modification of database contents or unauthorized access to the administrator account.

Protection against Cross-site scripting (XSS)

Cross-site scripting is a vulnerability where malicious users can include Javascript code into webpages, through entering a script in web forms, or including code in variables which are being printed on the web page. This can, for instance, be used to forward a user unnoticed to a different site. The protection against XSS is the reason most forms in LOVD do not allow HTML content.

Password protected data manipulation forms

To prevent manipulation of your variant or patient data when your computer is used by someone else while you are logged into LOVD, all data manipulation forms are password protected. Before the form can be submitted and processed, your password is required to confirm the changes. This also protects you against Cross-site request forgery (CSRF) attacks.
12.2 What you can do to protect your data

A chain is only as strong as the weakest link. Even with all the effort we have put into making LOVD as safe as possible, some weak links may still exist. Read this section to learn about how to protect your data in LOVD further.

1. Choose your password carefully and protect it. Even though LOVD does include a password quality check, please take care in choosing a password that can not be guessed by others and you don’t use for other systems as well. Also, don’t write it down on a sticky note stuck near your computer. In case you’re the system administrator who installed LOVD, make sure the MySQL password is also chosen securely.

2. Don’t leave your computer unattended while logged in. This will give anyone who uses your computer in your absence access to the data stored in LOVD. Also, don’t leave your computer unattended if you’re not logged in but had your browser remember your password for you. Once you’re done using LOVD, log out.

3. Be careful where you leave your backup files. Making backups of your data in LOVD is a smart thing to do. Errors made by both machine (hardware error) or man (accidentally deleting data) can be solved easily if you have proper backups. However, these backups do contain all your sensitive patient and variant data in a non-encrypted way. Be careful where you store your backup files. Don’t store them on network drives accessible by more people than the ones who you would allow to access LOVD itself.

4. Use SSL. Ask your systems administrator to enable SSL (https) on the server your LOVD is installed on, and enable the “Use SSL only to access LOVD” option in the LOVD system settings. This encrypts any data exchange between you and LOVD, such as your password and sensitive patient information.

5. Don’t allow access to your LOVD files or the database to other users on the server. If there are other users on the server you have LOVD hosted on, make sure they cannot access the LOVD directory or get access to the same database LOVD stores its information in.
13. Updating LOVD

Several times a year, a new LOVD build is released. An update may contain bug fixes, new features, new custom columns and such. Therefore, each update has a priority setting to indicate how important it is to update your LOVD installation to the new release.

Depending on whether or not LOVD has found that there is an update available, an update icon appears at the bottom right-hand side of the screen:

- There is an update available.
- There is no update available.
- There was an error while checking for updates.

If you are logged in as an authorized user, you can click the update notification icon for more information on the status of the update. If an update is available, detailed information is presented:

**Last checked for updates:** Shows the date LOVD last checked for a new update.

**Latest version:** Shows the detected update.

**Release date:** The date this update has been released.

**Priority level:** Shows how important the LOVD developers consider this update. Can contain “Optional”, “Common”, “Suggested”, “Recommended”, “Important” or “Critical”, with increasing priority.

**Release info:** Information on the changes that have been done in this release. Please note that this is usually only a very short list of updates. A more elaborate description can always be found in the news section of the LOVD website, and the full list can be found in the LOVD changelog.

If you choose to upgrade LOVD, follow these steps:

1. Download the new build. Either using the direct link to it from the update window, or from the website.
2. Unpack the new build and overwrite the LOVD files.
3. If you’re working from your local machine, upload the updated files to your webserver.
4. Log into LOVD. This is an important step. If you are already logged in, perform some activity, like clicking on the Setup tab. LOVD will then upgrade the database tables to make sure the database backend has the same version as the files. Skipping this step may result in serious errors occurring.
5. You’re done!
14. Troubleshooting

In case you run into any kind of problem or when you have a question, we recommend you first check the manual to see if perhaps the answer can be found there. A newer version of the manual may be available online, on the LOVD website. If that doesn’t resolve your problem or answer your question, please follow the following steps:

- Check the Frequently Asked Questions on our website to see if that provides you any answers.

- If not, check the LOVD forum to see if that helps you.

- If not, please submit your question to the LOVD3 support forum.

- If you’re sure the problem you are having is an LOVD bug, please use our bug tracking system to report your issue.

- Only in case you don’t want to publicly put your problem or question on the forum or bug tracking system, you can contact us also directly through the website.