

SpaceLight Command Line Documentation Version 1.5

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Contents

6	Further Reading, References	13				
5	Descriptor Choices	12				
4	Command Line Options4.1Overview4.2Minimal Required Options4.3Program Options4.4Configuration4.5General Options	5 5 6 9 10				
3	3 Jump Start: Finding Analogs in Large Chemical Spaces					
2	Technical Prerequisites	3				
1	Introduction	2				

1 Introduction

All links, references, table of contents lines etc. in this document are clickable.

Please note that this package is a command line package.

SpaceLight is a command line package to navigate combinatorial chemical spaces ("fragment spaces") of multi-billion size and beyond using classical similarity descriptors (fingerprints). SpaceLight is also part of our flagship platform infiniSee and operates in the background of the Analog Hunter mode (https://www.biosolveit.de/download/?product=infinisee).

SpaceLight is a perfect complement to our fuzzy pharmacophore-based FeatureTree descriptor (FTrees, https://www.biosolveit.de/download/?product=ftrees). Whereas the FeatureTree descriptor has a pronounced strength in detecting **distant** neighbors with chemical similarity, SpaceLight will find the **close-by** neighbors in a chemical space.

SpaceLight lets you

- conduct very fast, **fingerprint-based similarity searching** across vast combinatorial Chemical Spaces [3] for your query molecules
- perform similarity searches using a variety of different fingerprint types and sizes (ECFP and CSFPs, Section 5)
- perform similarity searches in enumerated library files (SMILES, SDF or MOL2)
- visualize the local similarities of the underlying fragments from which the result molecule is constructed (Figure 2)

SpaceLight traverses huge chemical spaces using Lego-like chemical reaction combinatorics behind the scenes. To conduct quick calculations, we formalize reactions and encode them as pseudo-linking reactions. Per reaction, one so-called "topology graph" is stored and every node of this graph contains formalized, virtual building blocks, or as we call them, "reaction fate foreseeing" fragments (Figure 1). For more information on fragment space generation, please have a look at our CoLibri package (https://www.biosolveit.de/products/#CoLibri).

The SpaceLight search uses fast combinatorial algorithms and can deliver results in a few seconds only — even for very large, multi-billion sized spaces and beyond. The fingerprint-based Tanimoto similarities are calculated on the fragments, leading to local similarities at first. With the combinatorics behind every topology graph, the most similar hit molecules can be constructed and enumerated from the fragments in a very time-efficient manner. Detailed information on the basic ideas of the algorithm can be found in the original publication by Bellmann et al.[2]

The implemented 2D similarity descriptors (CSFPs) are specially optimized for similarity searches in combinatorial fragment spaces (Section 5).[1] They are automatically prepared for every chemical space during space generation with the CoLibri package (SpaceLightDBCreator, see link above). Additionally, the extended connectivity fingerprint (ECFP) known from 2D similarity searches in traditional libraries can also be used. Spaces ready to be searched with SpaceLight can be downloaded from our website (https://www.biosolveit.de/chemical-spaces/).

Summarizing, with SpaceLight you can conduct 2D similarity searches with well-known fingerprints (e.g. ECFP4) in much bigger spaces than with other methods, and you also require much less time, making it possible to search vast spaces even on modest, standard hardware.



Figure 1: Example of topology graph representing a reaction with combinatorial fragments.

2 Technical Prerequisites

SpaceLight is a command line application. It needs the following to run:

• The SpaceLight package

(https://biosolveit.de/download/?product=spacelight)
Depending on your operating system, some libraries may have to be installed (get in touch with
us: mailto:support@biosolveit.com; and please mention any errors/warnings that you see in
your mail)

- A shell (Linux/Unix) or a terminal (macOS), or a command line environment (Windows; e.g.: cmd.exe)
- A valid license (from mailto:license@biosolveit.com)

The license setup instructions will come with the license that we will send out — or has already been sent out to you. A "test license" that you can request online and that is sent to you instantaneously can simply be placed next to the executable (spacelight.exe, spacelight, or SpaceLight — depending on your operating system). For macos please read on...

macOS Specialties On macOS, the executable will typically reside inside the *.app package:

/Applications/SpaceLight.app/Contents/MacOS/SpaceLight

To place the short term test license there, you will have to go into the *.app package using a right mouse click on SpaceLight.app in the Finder, and click on "Show package contents". In there, you will see the Contents/ subfolder, in there the MacOS subfolder, and in there, the SpaceLight executable. If you are about to use the **test license**, place it right there, next to the executable. A longer term license will be handled separately, we will tell you how when we send that very license.

When you call SpaceLight for the first time, go to the Finder, and navigate to the Applications folder. Do a right(!) click on SpaceLight.app, and — if applicable — confirm that you want to open the program. It will flash up once, and you are good to go at the terminal prompt from there on.

To make the first step, call SpaceLight within your shell/terminal/environment.

3 Jump Start: Finding Analogs in Large Chemical Spaces

Your license is all set? You unpacked the installation archive? You downloaded or prepared a fragment space file (.space file from https://www.biosolveit.de/chemical-spaces/or .tfsdb file)? Then here is a typical call to search a query against a fragment space:

```
./spacelight -i <path/query.sdf> -s <path/fragmentspace.space>
```

./spacelight -i <path/query.sdf> -s <path/fragmentspace.tfsdb>

The query can be an SD file (.sdf), a SMILES file (.smi or .smiles, containing line-separated SMILES), a .mol or .mol2 file with one or multiple entries. For quick searches, the input can just as well be a SMILES string enclosed by quotation marks, for example:

```
./spacelight -i "CC(C)C(=0)N" -s <path/fragmentspace.space>
```

By default, the 100 most similar hit molecules are written as SMILES to your console/shell (STDOUT). To write your results to an output file (.csv or .sdf), additionally use either the -o / --output-files option (writes a separate output file for every query) or the -O / --single-output-files option (writes a single output file with the concatenated results for all queries), for example:

```
./spacelight -i "CC(C)C(=0)N" -s <path/fragmentspace.space> -o <path/output.csv>
```

The output file contains the structure of the result molecules as well as detailed information on the used fingerprint type and similarity score (see page 7 for more information). You can adjust the number of results by using the --max-nof-solutions option:

The fragment connected subgraph fingerprint (fCSFP4) is used as the default similarity descriptor (see page 9).[1] The Tanimoto similarity is used as similarity measure. The similarity values are normalized between 0 (no similarity) and 1 (identical, in the framework of the descriptor). The default descriptor is **not** stereo-aware, but it captures elements, connectivity, valence, and ring membership. As such it will find "near-neighbor" similar compounds in a chemical space. You can limit the output to those results which exceed a certain similarity by using the **--min-similarity-threshold** option:

Instead of CSFP fingerprints, you can also use the well-known extended connectivity fingerprints (ECFP, see page 9 and Section 5):

```
./spacelight -i "CC(C)C(=0)N" -s <path/fragmentspace.space> -f ecfp4
```

4 Command Line Options

4.1 Overview

An overview of all command line options is available by calling SpaceLight with --help.

```
./spacelight --help
Program options:
-i [ --input ] arg
                                         Input query molecule file or single input molecule as smiles.
                                         Supported file types are *.smi, *.smiles, *.mol, *.mol2 and *.sdf.
Paths to library input molecule files for similarity scoring or to
-s [ --search-files ] arg
                                         Topological Fragment Space database files or Fragment Spaces
                                         Supported file types are *.smi, *.smiles, *.mol, *.mol2, *.sdf,
                                         *.tfsdb, *.space and *.zip.
                                         Output base files (suffixes are required). For each query molecule,
-o [ --output-files ] arg
                                         the results are written to a separate output file. Supported file
                                         types are *.csv and *.sdf.
-O [ --single-output-files ] arg
                                         Output files (suffixes are required). All results are written to a
                                         single output file. Supported file types are *.csv and *.sdf.
                                         Output base file name for matching images (suffix required).
Supported file types are *.pdf, *.png and *.svg.
-m [ --match-image-base-file ] arg
                                         Note: For each match a separate file is created.
--gen-mapping-output [=arg(=1)]
                                         Generates command line SpaceLight similarity descriptors and annotates
                                         them in the output file.
Configuration:
                                         Fingerprints for searching: ECFP and CSFP variants are supported. Supported ECFP variants are ECFP0 to ECFP8. Three CSFP subtypes are
-f [ --fingerprint ] arg (=fCSFP4)
                                         available: fCSFP, iCSFP and tCSFP in variants from 1 to 5. E.g. ECFP4,
                                         fCSFP5, iCSFP4 or tCSFP3.
--min-similarity-threshold arg (=0) Similarity threshold below which molecules are discarded [0.0 to 1.0].
--max-nof-results arg (=100)
                                         Maximum number of top-ranking result molecules [1 to 1000000].
--expand-alternative-results [=arg(=1)]
                                         Write alternative results based on alternative reaction paths.
General options:
-h [ --help ]
                                         Print this help message
--license-info
                                         Print license info
--thread-count arg
                                         Maximum number of threads used for calculations. The default is to use
                                         all available logical cores.
                                         Print version info
--version
-v [ --verbosity ] arg (=2)
                                         Set verbosity level
                                               0 [silent]
                                               1 [error]
                                               2 [warning]
                                               3 [workflow]
                                               4 [steps]
```

The abbreviated, one-letter options are preceded with one dash – whereas the longer, named options are preceded with two dashes: --. If an option needs an argument (arg), you can include or omit the equals sign. Adapt the command line usage to your operating system and shell.

4.2 Minimal Required Options

This section describes the arguments you must specify at minimum to successfully run a similarity search. First, you must provide the path to a file containing the query compounds. All well-known data formats are supported (MOL, SDF, SMILES, MOL2). Instead of a file containing the query molecules you can also specify a single SMILES string enclosed by quotation marks. The SMILES string or the molecule file must be passed to the -i option. Additionally, you need to specify the path to a fragment space (.space file) to be searched. Alternatively, you can also specify a library file (SDF, MOL2, SMILES file) to perform an enumerated search instead of a space search. Either the space file or library file have to be specified with the -s option. The minimal search prompt then has the following general form:

```
<path/to/spacelight/executable> -i <path/to/queries> -s <path/to/space_file>
```

When you specify the required paths the search prompt might look like the following:

./spacelight -i my_queries.sdf -s my_space.space

Or, if you specified a SMILES string instead of a file:

./spacelight -i "CC(C)C(=O)N" -s my_space.space

In the examples above, the output is printed on the console by default, which will look similar to the following example:

Query: O(CCCC)C O(CCCC)C Rank: 1 sim: 0.706 O(CCCDC)C EN300-1717039 Rank: 2 sim: 0.577 O(CCC[NH2+]CCCC)C m_270004cba___8288582___9143638 Rank: 3 sim: 0.577 S(CCCDC)CCCC m_62bba___875776___9108904 Rank: 4 sim: 0.536 O(CCCC[NH2+]CCCC)C m_270004cba___8290272___9143638 Rank: 5 sim: 0.536 S(CCCCDC)CCCC m_62bba___875776___10159086 ...

For every query molecule, the respective result molecules are printed to the console with information on the rank, similarity score, SMILES representation and name of the molecule. By default, 100 results molecules per query are printed to the console. If you want to increase or decrease that number, please have a look at the **--max-nof-results** option (see page 10).

Of course, you can also write the result molecules to an output file either with the -o option (one separate output file per query) or with the -0 option (a single output file with results molecules for all queries). See page 7 for more information.

4.3 Program Options

-i [--input] arg Specify a file containing the query molecules. Supported file formats are SDF, MOL and MOL2. You can also provide a text file containing multiple line-separated SMILES (file extension must be .smi or .smiles). Instead of a query file you can also specify a single SMILES string enclosed by quotation marks. It is also possible to use the -i option multiple times in a row (see examples below). NOTE: The -i option is required.

Examples:

spacelight -i myquery.sdf
spacelight -i myquery.sdf -i mydrugs.smi
spacelight -i "CC1=CC=CN=C1"

-s[--search-files] arg Specify a topological fragment space file (.tfsdb) or a library file (.sdf, .mol2, .smi, .smiles) or a space file (.space). This file is searched for close analogs to the query molecules given with the -i option. You can also search multiple spaces or library files at once by using the -s option several times in a row. And even libraries and chemical spaces can be mixed (see examples below).

NOTE: The -s option is required.

Examples:

spacelight -s mylibrary.sdf
spacelight -s mylibrary.sdf -s myspace.space
spacelight -s space1.space -s space2.space

-o [--output-files] arg Specify the base name for the output files as argument here. The output will be written either as SD file (.sdf) or .csv file — or both. Specify the desired output file type by the file extension. The SD file will contain additional information for every result molecule in dedicated SD data fields (see below). The CSV file will contain the result molecules as SMILES together with additional information (see below).

NOTE: If you have multiple queries in your input file, then a separate CSV or SDF file will be written per query! To write all results from multi-query input files in a single output file, see **--single-output-files** option below.

NOTE: If you do not specify an output file, reduced output will be written to the command line (STDOUT, see Section 4.2).

Examples:

spacelight -o myoutput.sdf

spacelight -o myoutputtable.csv

```
spacelight -o myoutput.sdf myoutputtable.csv
```

The latter example outputs both, one SD file and one CSV file per query contained in your input file. The names of the output files will have the following general structure:

myoutput_{querynumber}.sdf and myoutputtable_{querynumber}.csv

The output file(s) contain additional information for every result molecule:

- **result rank**: rank among all results for this particular query
- **fingerprint**: fingerprint descriptor used for the search (see Section 5)
- **fingerprint similarity**: Tanimoto similarity value (derived from query molecule fingerprint versus result molecule fingerprint)
- result name: name of the result molecule
- query name: name of the query molecule
- **query smiles**: SMILES of the query molecule
- **space**: name of the searched space(s) or library
- reaction name: name of the reaction that constructs the result molecule
- **reagent(1-5) name**: name of building block (1-5) from which the result molecule is constructed (relevant only for space searches, number depends on reaction)
- **reagent(1-5) smiles**: SMILES of building block (1-5) from which the result molecule is constructed (relevant only for space searches, number depends on reaction)



Figure 2: Example of a match image. The query is on the left side, the hit molecule on the right side.

-O [--single-output-files] arg Specify the name for the output files as argument. The output will be written either as SD file (.sdf) or .csv file — or both. Specify the desired output file type by the file extension. The SD file will contain additional information for every result molecule in dedicated SD data fields (see below). The CSV file will contain the result molecules as SMILES together with additional information (see below).

As a difference to the --output-files option (see above), all results from multi-query input files will be written to a single output file (concatenated results). It is also possible to write both a CSV file and a SD file at the same time (see last example below).

NOTE: If you do not specify an output file, reduced output will be written to the command line (STDOUT, see Section 4.2).

Examples:

spacelight -0 singleoutput.sdf

spacelight -0 singleoutput.csv

spacelight -0 singleoutput.sdf singleoutput.csv

-m [--match-image-base-file] arg Specify a base name for the match image output files as argument. This will generate (one per hit molecule, so potentially many!) output images that explain the matching of query versus hit molecule in 2D pictures. Depending on the file extension you specified, images will be written as .png, .pdf, or vector-based .svg files. NOTE: Generation of match images leads to extended runtimes.

Example:

spacelight -m matching.png

This call will generate one .png file per hit molecule(!), the file names will look like:

matching_{querynumber}_{hitnumber}.png.

Figure 2 shows an example matching image. The local similarities are listed and color-coded on the right hand side. The query molecule is located on the left side (Tofacinib), the hit molecule on the right side. The hit molecule is colored according to the fragments/building blocks from which it is constructed. As you can see, the aromatic heterocyclic system (cyan) is identical (similarity 1) between query and hit, the part with aliphatic ring (orange) is less similar (similarity 0.507).

--gen-mapping-output If you specify this option, for every hit molecule the local similarity values (annotated in the *similarity-descriptor* column/tag) and the corresponding substructures (annotated as semicolon-separated SMILES in the *similarity-descriptor-smiles* column/tag) are written to the output files. This is the same information that is visualized for the hit molecule in a match image (see Figure 2 and the description for the --match-image-base-file option above). NOTE: Calculation of local/fragment similarities increases the runtime!

Example:

spacelight --gen-mapping-output

4.4 Configuration

With the options described in this section you can adjust the algorithmic parameters of SpaceLight. For example, you can modify the fingerprint used to derive similarity, adjust the minimal similarity threshold below which result molecules are discarded and change the maximum number of results generated for each query molecule. All parameters in this section have default values which are round-bracketed in the following.

-f [**--fingerprint**] **arg**(**=fCSFP4**) With this option, you can adjust the fingerprint type and size used for the determination of similarity between query and result molecules. For more information on the available descriptors see Section 5. Shortly, you can choose between three different versions of the connected subgraph fingerprint (CSFP [1]), each version with features containing from 1 up to a maximum of 5 heavy atoms (CSFP1-5): fCSFP1, fCSFP2, fCSFP3, fCSFP4 (default), fCSFP5; iCSFP1, iCSFP2, iCSFP3, iCSFP4, iCSFP5; tCSFP1, tCSFP2, tCSFP3, tCSFP4, tCSFP5. Additionally, you can choose the "classical" circular extended connectivity fingerprint (ECFP [4]) with a maximum diameter of 8: ECFP0, ECFP2, ECFP4, ECFP6, ECFP8.

NOTE: Specification is case-insensitive.

Examples:

spacelight -f ecfp4

spacelight -f fcsfp5

spacelight -f tCSFP3

--min-similarity-threshold arg(=0) Takes a number between 0 and 1 as argument. This parameter adjusts the minimum similarity threshold below which the result molecules are discarded. By default, the value is 0, e.g. no result molecules are discarded.

Example:

spacelight --min-similarity-threshold 0.7

--max-nof-results arg(=100) Takes a number between 1 and 1,000,000 as argument. You can adjust the maximum number of result molecules per query which will be written to the output files. The default value is a maximum of 100 results per query. The results will always be sorted by descending Tanimoto similarity, so the parameter controls the TOP number of results. The output is limited to 1,000,000 result molecules.

Example:

spacelight --max-nof-results 5000

--expand-alternative-results Expands alternative reactions for a hit molecule. In chemical spaces, the same molecule can be formed in different reactions with different reagents/building blocks. If you use this option, these different possibilities are written to the output. Alternative results all have the same rank and the same similarity score but have different names and different reagents. You can also use this option to find identical results in different spaces if you specify multiple spaces at the same time (see -s option).

Example:

```
spacelight --expand-alternative-results
```

4.5 General Options

-h[**--help**] Displays the command line help with short descriptions for every argument option. For more information see Section 4.1.

Example:

spacelight --help

--license-info Shows command line information about the license setup you currently use. If you have any problems with your license, send an email to mailto:support@biosolveit.com and include this information.

Example:

spacelight --license-info

--thread-count arg Specify the maximum number of threads used for your similarity searches. By default, all available logical cores of your computer are used. You may want to reduce the number of threads if you want to run other computations on your computer at the same time, or if you share the compute resource.

Example:

spacelight --thread-count 4

--version Displays information on the version of SpaceLight on the command line. In quoting Space-Light, please mention this version number.

Example:

spacelight --version

-v [**--verbosity**] **arg(=2)** Set the verbosity level, e.g., the level of console output, with an integer argument. The default value is 2. The following options are available:

- **o** Silent. No messages will be displayed in the console during the similarity search run. Errors will be ignored whenever possible.
- 1 Error. Only error messages will be displayed.
- 2 Warning. The default setting, warnings and error messages will be displayed.
- **3** Workflow. In addition to errors and warnings, information on the different steps of the similarity search are displayed on the command line.
- 4 Steps. In addition to the 'Workflow' option, the progress of each step is displayed in detail.

Example:

spacelight -v 0

5 Descriptor Choices

SpaceLight has several built-in similarity descriptors (fingerprints, see page 9). Depending on your use case, one descriptor may be more suited than another. The acronym CSFP stands for *Connected Subgraph Fingerprints*, they describe all possible chemical features ("substructures") containing up to 5 heavy atoms for a given molecule.[1] In contrast, the "classical" ECFP descriptors (*Extended Connectivity Fingerprints*) describe molecules with a circular collection of features ("sit on one atom, then collect sphere with radius 1 around you, then sphere with radius 2, and so on").[4] CSFP descriptors are particularly optimized for similarity searches in fragment spaces as their set of features minimizes information loss across fragment boundaries. See the table below that has been taken from the original SpaceLight publication; it may serve as an overview of the respective atomic properties stored for the different descriptors along with the respective features.[2]

		ECFP	fCSFP	iCSFP	tCSFP
chemical substructures	circular	х			
	all		x	x	x
	element	x	x	х	x
	connectivity	x	x		x
	connectivity in substructure		х	x	
	valence	х	x	х	
atom properties	valence in substructure		х	x	
	aromaticity				x
	π electrons				
	formal charge	x			
	weight	x			
	ring membership	X	x		

Summarizing, we suggest to use the following descriptors for the respective use cases:

- fCSFP (fragment CSFP): Use this descriptor when the results should be **highly similar** to the query compounds. This is likely the descriptor that you should be able to relate most to when comparing it with the traditional ECFP descriptors. Element, valence state, and connectivity (within the feature and to the surroundings of the feature) are captured for every atom.
- tCSFP (topological CSFP): Compared to the fCSFP, tCSFP does not take into account the atom connectivity within substructures (features) nor the valence states, but on the other hand information on aromaticity is stored. This makes the tCSFP less strict in deriving similarity for a given compound pair compared to the fCSFP. Use this fingerprint in cases you do not find close analogs with fCSFP or ECFP fingerprints.
- iCSFP (independent CSFP): This descriptor is recommended for **substructure retrieval**. The iCSFP describes all structural features of a compound by properties of the atoms that are *independent* of the surroundings of its substructures. Therefore, it is suitable when searching for molecules that contain substructures or very similar substructures of a given query compound in an arbitrary order.
- ECFP: Extended Connectivity Fingerprints are widely used in traditional similarity searching. Use ECFPs when you would like to stay close to or compare to another tool that uses this descriptor type.

6 Further Reading, References

The original ideas behind the SpaceLight method are covered in the publication by Louis Bellmann and Matthias Rarey.[2]

If you prefer to control the SpaceLight algorithm through a graphical interface, download our platform infiniSee. Similarity searches with ECFP4 fingerprint can be performed in the Analog Hunter mode. (https://www.biosolveit.de/download/?product=infinisee)

References

- [1] Louis Bellmann, Patrick Penner, and Matthias Rarey. Connected subgraph fingerprints: Representing molecules using exhaustive subgraph enumeration. *J. Chem. Inf. Model.*, 59(11):4625–4635, 2019.
- [2] Louis Bellmann, Patrick Penner, and Matthias Rarey. Topological Similarity Search in Large Combinatorial Fragment Spaces. J. Chem. Inf. Model., 61(1):238–251, 2021.
- [3] Torsten Hoffmann and Marcus Gastreich. The next level in chemical space navigation: going far beyond enumerable compound libraries. *Drug Discovery Today*, 24(5):1148–1156, 2019.
- [4] David Rogers and Mathew Hahn. Extended-connectivity fingerprints. J. Chem. Inf. Model., 50(5):742–754, 2010.

We wish you great success and much joy with SpaceLight!