Parent-map

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Overview

Parent-map analyzes protein or DNA sequences which are derived from one or multiple parent sequences, and shows parental contributions as well as differences from relevant parents.

Originally developed to analyze capsid protein sequences obtained by directed evolution, Parent-map can be used in any case where variant sequences are to be compared to parent sequences from which they are derived. Parent-map detects sequence shuffling as well as substitutions, insertions and deletions.

Parent-map takes two FASTA files as input and generates four text files and one html file as output.

The first input file contains the variant sequences to be analyzed, in either single- or multi-line FASTA format. Sequences are read into memory one at a time, so there is no limit to their number. They don't need to be related, and can be derived from any combination of parental libraries.

The other input file contains the parental sequences. FASTA format is only required if more than one parent is present. All parental sequences are loaded into memory together, as their number is limited.

The output files have the following endings in their name: -stats.txt (statistics), -par.txt (parental contributions), - par.html (parental contributions, colorized html file), -def.txt (sequence definitions) and -aln.txt (alignments).

The -stats.txt file is a simple table showing an overview of the variant sequences main features. Sequences are sorted by the number of contributing parents, then by the parent name.

The -par.txt file is an alignment of each variant against all its contributing parents. This file is particularly useful for variants derived from shuffled libraries, as it maps the most likely parental contributions. Contributing parental fragments are displayed using an identity character (which can be customized). Obviously, in the case of multiple parents, the combination of fragments shown is only one among a very large number of possible combinations. However, the solution shown has the largest fragment lengths, the smallest number of fragments and the smallest number of parents.

The -par.html file is a colorized version of the -par.txt file, with each parent represented with a different color.

The -def.txt file gives a formal definition of each variant sequence in relation to its parent(s), in a table format. It's a blueprint allowing to reconstruct the variant sequences from a minimal set of instructions.

The -aln file aligns variants under their main parent. As opposed to the -par.txt file where there is an alignment per variant, here there is an alignment per parent. First, alignments for single parent sequences are shown, then alignments for multi-parent sequences if they exist, against their main parent. For legibility, only regions in which at least one

of the variants differ from the parent are shown. Each region is numbered according to its position in the parental sequence.

Parent-map is a platform-independent Python 3 script available as a bioconda package, as well as a Windows executable. It can be run in command-line mode inside a terminal or in a graphic user interface. Starting it without arguments will launch the graphic interface, while adding any argument will run it as a command-line program. The Windows version can also do both, with the only difference that it will not display anything in command-line mode.

Source code: https://github.com/damienmarsic/Parent-map

Python package: https://pypi.org/project/parent-map/

Bioconda recipe and package: http://bioconda.github.io/recipes/parent-map/README.html

Windows executable: https://sourceforge.net/projects/parent-map/

How to cite

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BibTeX entry: @article{Marsic2021, doi = {10.21105/joss.02864}, url = {https://doi.org/10.21105/joss.02864}, year = {2021}, publisher = {The Open Journal}, volume = {6}, number = {57}, pages = {2864}, author = {Damien Marsic}, title = {Parent-map: analysis of parental contributions to evolved or engineered protein or DNA sequences}, journal = {Journal of Open Source Software} }

Installation

The recommended method is to install parent-map as a bioconda package under a Python 3 conda environment. If you don't have such environment, first install Miniconda3 from here (exists for Linux, MacOSX and Windows): https://docs.conda.io/en/latest/miniconda.html

To install the parent-map bioconda package, at the conda prompt type:

conda install -c bioconda parent-map

Windows users who prefer not to use a conda environment can install the Windows executable version of parent-map from here: https://sourceforge.net/projects/parent-map/

Parent-map can also be installed using pip:

pip install parent-map

However, dependencies might need to be installed individually. If you get an error message about missing modules when trying to run parent-map after installing it with pip, use pip install to install missing modules. Example:

pip install gooey

Usage

To run parent-map from a Python 3 conda environment, type:

python -m parent-map

This will launch the graphic interface (same as if launching the Windows version).

To run in command-line mode, add arguments. For example:

```
python -m parent-map -v python -m parent-map -h
```

In the first example, the version will be displayed. The second example will show a help message.

Typical usage:

python -m parent-map variant_file parent_file optional_arguments

where variant_file is the name of the file containing the variant sequences, parent_file is the name of the file containing the parental sequences, and optional_arguments can be any combination of the optional arguments described in detail below.

Real-life example:

python -m parent-map example_variants.fasta example_parents.fasta -o test

This will result in the following output:

```
Sequence name 'AAV2-sept-Y-F' replaced with 'S16'
Parental maps saved into file: test-par.txt
Stats saved into file: test-stats.txt
Sequence definitions saved into file: test-def.txt
Alignments saved into file: test-aln.txt
Colorized version of parental map file saved into file: test-par.html
```

Note that the prefix 'test' entered with the -o (output) argument became the prefix of the 4 output files. Also note that the sequence name change (first line) can be avoided by using the -s argument (maximum name size) followed by the value 13 (the sequence name 'AAV2-sept-Y-F' is 13 character long, while the default value for -s is 12).

Parent-map can also bypass the sequence analysis and just colorize an existing -par.txt file into a -par.html file. This is useful when the user is not satisfied with the default colors and wants to try different color schemes. In this case, the -C argument is used followed by the name of the -par.txt file. Also, the -p argument must be used followed by a list of parent color pairs. Example:

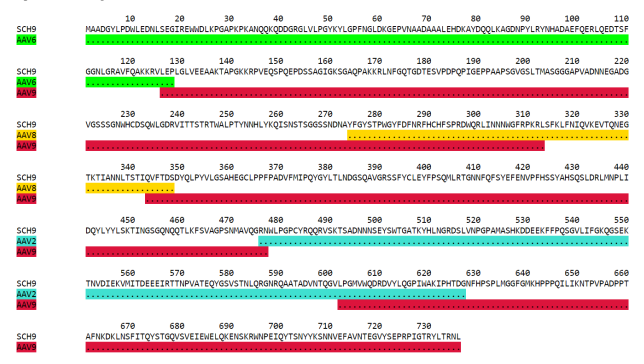
```
python -m parent-map -d -C test-par.txt -p 'AAV2 turquoise, AAV3B tomato, AAV5_

→lavender, AAV8 gold, AAV9 crimson, AAV6 lime'
```

The following output will be generated, while the newly created html file is automatically displayed (thanks to option -d) in the default web browser:

Colorized version of parental map file saved into file: test-par.html

A portion of the generated html file is shown here:



4.1 Arguments

The same parameters are available in both modes (graphic and command-line) but some default values might differ.

• Version: -v

Whenever the argument -v is used, parent-map will only display the program version and exit, no matter whether other arguments are present or not. In graphic mode, version information is under 'About' in the main menu.

• Help: -h

Whenever the -h argument is used, parent-map will only display the help message and exit, no matter whether other arguments are present or not (except -v). In graphic mode, the 'Help' menu item links to the documentation web page.

• Variants: file name or path to file name

The variant file must be in FASTA format, no matter how many sequences are present. In graphic mode, a Browse button allows to select the file from any directory. If a blank space is present within a directory or file name, the whole path needs to be surrounded by quotation marks, even in graphic mode. For example: 'D:my foldermy variants.fasta'

• Parents: file name or path to file name

The parent file must be in FASTA format unless a single parent is present, in which case raw sequence (with no name) is fine. Again, a Browse button is there in graphic mode. If a blank space is present within a directory or file name, the whole path needs to be surrounded by quotation marks, even in graphic mode. For example: 'D:my foldermy parent.fasta'. It is possible to change the position numbering by adding a number after the sequence name (separated by a blank space). That number will then become the position number of the first character in the sequence. If no number is provided, the first character will be considered to be in position 1. This is useful when standard numberings need to be used but only a region of the gene is sequenced.

• Output: -o

The output files will be saved in the same directory as the variant file. If this argument is used, the prefix entered will be the part of the file names before the -stats.tx, -par.txt, -def.txt and -aln.txt endings. If this argument is not used, a prefix will be automatically generated from the variants and parents file names, as well as some arguments and their values.

• MinFragLen: -m

This is the minimum fragment length used to map variant regions against parental sequences. Default values are 6 for protein sequences and 18 for DNA sequences (sequences with non [A, T, G, C] characters are automatically defined as 'protein'). It is recommended to keep the default values.

• MinOverlap: -V

This is the minimum overlap length in the case of sequences produced by shuffling parental sequences. Default values are 2 for protein and 6 for DNA. It can be useful to decrease the value in cases where variants from multiple parents are not produced by overlapping randomly digested fragments, but rather by rationally designed combinations.

• MaxNameSize: -s

Maximum length of sequence names before the name is replaced with a shorter generic name ('S' followed by a number corresponding to the sequence position in the FASTA file). Default value is 12.

• SeqChars: -c

Number of sequence characters per line in alignments (-par.txt and -aln.txt files). Default value is 120.

• LowerCase: -1

By default, all sequences are displayed in upper case. Use the -l argument to display sequences in lower case instead.

• VRSides: -e

Number of characters to be included each side of variable regions in the -aln.txt file. Note that increasing the value results in neighboring regions merging. Modifying this value only alters the esthetics of the displayed alignments. Default value is 1.

• Overwrite: -f

By default, if an output file with the same name already exists, parent-map exits with a warning to either change the output prefix or to use the -f argument. The -f argument forces overwriting of existing files with identical names without warning.

• Symbols: -S

Symbols for identity (default: '.') and gap (default: '-') in -par.txt and -aln.txt files. Symbols should be exactly 2 non identical, non alphabetical and non blank characters.

• DisplayResults: -d

The output files are automatically displayed in a web browser by default in graphic mode only. Use the -d argument to automatically display in command-line mode. The option might not work, depending on the user's operating system and browser configuration. The txt files might be displayed in the default application for text files instead, if the browser is unresponsive, or might not display at all. In any case, the 5 output files are always saved in the same directory as the input files, and can be opened at any time using any appropriate displaying application (a web browser is recommended). Note that if a word processor is used, a monospaced font (such as Courier New for example) needs to be applied to the contents for proper display. Note that even when using a monospaced font, Microsoft Word uses shorter blank spaces, which distorts the display. In this case, replace all blank spaces in the document with a monospaced blank space created by pressing the space bar while simultaneously holding both Shift and Ctrl keys.

• Colors: -p

When parent-map is used to analyze sequences (using the 2 input files), the -p (palette) argument is unnecessary (default colors are used) unless the user wishes to apply a specific color scheme, in which case -p must be followed by a list of parent color pairs in the following format (pairs separated by commas, blank space between parent and color, list surrounded by single or double quotes):

-p 'parent1 color1, parent2 color2, parent2 color3,...'

Parent names must be exactly as they appear in the -par.txt file, including the case. Color names are case-insensitive and must be standard html colors, such as shown on this page: http://www.html-color-names.com/color-chart.php

When sequence analysis is bypassed and parent-map is only used to colorize an existing -par.txt file (using the -C argument followed by the file name), the -p argument is required.

• ColorizeOnly: -C

This option is to bypass the sequence analysis and to only colorize an existing -par.txt file. The file name (or path) needs to be entered after -C.

Output files

5.1 Statistics

The -stats.txt file shows a summary of the variant sequences main features, in table format. Sequences are sorted by number of parental sequences, then by main parent name, then by the number of insertion sites. Description of the table columns:

- 1. **Name:** Variant sequence name as appearing in the variant FASTA file, or as a shorter name made of 'S' followed by the sequence number in the original file in case the original name is longer than the default maximum name length.
- 2. Length: Sequence lengths (number of amino acid residues for protein sequences, number of nucleotides for DNA sequences).
- 3. Parents: Number of parental sequences detected as having contributed to the variant sequence.
- 4. **Main:** Name of the main contributing parental sequence (the parent covering the largest area of the variant, in case of multiple parents).
- 5. **Coverage:** % of the variant sequence covered by the main parent. Note that this is not the same as global similarity to the main parent sequence in the case of multiple parents. In that case, it's about the parts of the variant sequence that are not already covered by other parents. It's actually the % of positions identical to the main parent as displayed in the -par.txt file.
- 6. Matches: Number of sequence positions matched (as opposed to a % as in the previous column).
- 7. **ID%:** Global similarity of the variant sequence compared to the main parent sequence (% of identical positions). In the case of a single parent, this is identical to coverage (6th column). However, in the case of multiple parents, this is the % identity with the parent along the whole variant sequence (and not just the part covered by that parent and not by other parents as in column 6).
- 8. **Identities:** Global similarity as the number of identical variant sequence positions compared with the main parent.
- 9. Ins_sites: Number of sites in the variant sequence that have an insertion compared with the main parent.
- 10. Ins: Total number of nucleotide or amino acid insertions in the whole variant sequence.

- 11. Del_sites: Number of sites in the variant sequence that have a deletion compared with the main parent.
- 12. Dels: Total number of nucleotide or amino acid deletions in the whole variant sequence.
- 13. Subs: Total number of substitutions in the variant sequence compared with the main parent.
- 14. **Other:** Number of sequence positions that could not be matched to any parent and could not be considered as a variation of one particular parent.

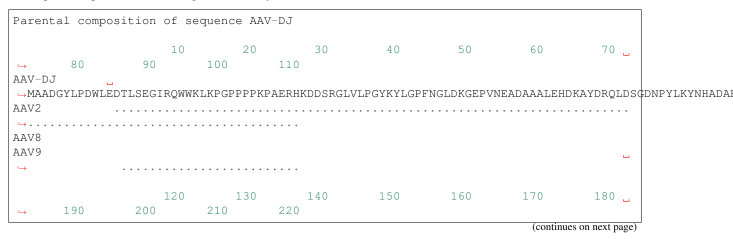
Example of a -stats.txt file:

	Length	Parents	Main	Coverage	Matches	ID%	Identities	Ins sites	Ins	Del sites	Dels	Subs	Other
Name	-			_				—		—			
LI-A	735	1	AAV2	98.10	721	98.10	721	0	0	0	0	14	0
LI-C	735	1	AAV2	99.46	731	99.46	731	0	0	0	0	4	0
AAV2-sept-Y-F	735	1	AAV2	99.05	728	99.05	728	0	0	0	0	7	0
7m8	745	1	AAV2	98.39	733	98.39	733	1	10	0	0	2	0
rAAV2-retro	745	1	AAV2	98.39	733	98.39	733	1	10	0	0	2	0
AAV-F	743	1	AAV9	99.06	736	99.06	736	1	7	0	0	0	0
AAV-PHP.B	743	1	AAV9	99.06	736	99.06	736	1	7	0	0	0	0
AAV2i8	735	2	AAV2	99.18	729	99.32	730	0	0	0	0	0	0
AAV2.5T	725	2	AAV5	83.86	608	94.21	683	0	0	0	0	1	0
AAV-DJ	737	3	AAV2	55.50	409	92.94	685	0	0	0	0	1	0
NP84	735	3	AAV2	84.90	624	98.50	724	0	0	0	0	2	0
OLIG001	737	4	AAV8	66.49	490	95.12	701	0	0	0	0	1	1
SCH2	737	4	AAV9	53.73	396	89.55	660	0	0	0	0	0	0
SCH9	736	4	AAV9	62.23	458	92.12	678	0	0	0	0	0	0

5.2 Parental mapping

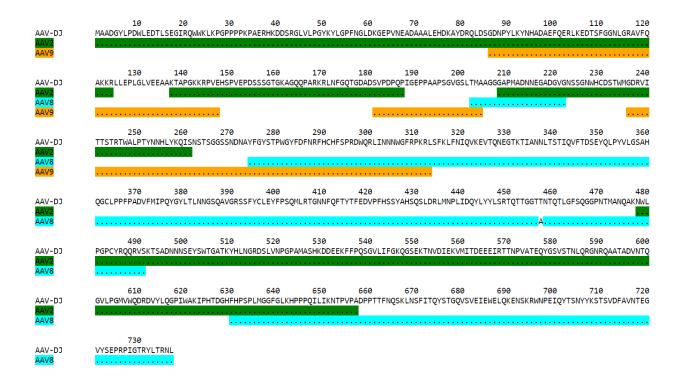
In the -par.txt file, each variant complete sequence is displayed along its contributing parental fragments. The parental mapping is simplified as much as possible, so that the variant is completely described using as few parents as possible and as few fragments as possible. Only the necessary parents are shown. If a fragment can be mapped equally to more than one parent, the parent that has the highest overall coverage is chosen. Identities are displayed in the parent lines using an identity symbol ('.' by default). If a mismatch occurs between 2 fragments mapped to the same parents or at a sequence end, it is considered a variation of the parent next to it and displayed as the parent sequence (substitution) or using the gap symbol ('-' by default) in the parent line for insertions or in the variant line for deletions. The -par.txt file is most useful for variant sequences selected from shuffled libraries. Two files are generated: a black and white text file (name ending with -par.txt) and a colorized html version of the same file (name ending with -par.html). In the html file, each parent name and its corresponding identity fragments are colorized using a different color (whenever possible). The default color scheme uses 12 different colors. If the number of parents in the file is greater than 12, the same set of 12 colors is used on the 12 next parents and so on. If the user wishes to apply specific colors, the *-p argument* needs to be used followed by a list of parent color pairs, either at the time of the sequence analysis or later with the -C argument to specify the -par.txt file to be colorized again.

Example of a -par.txt file (showing AAV-DJ only):



						(continued from	n previous page)	
AAV-DJ ⇔GGNLGRAVFQA AAV2	AKKRLLEPLGLVE		RPVEHSPVEPDS	SSGTGKAGQQ	PARKRLNFGQ	TGDADSVPDP	QPIGEPPAAPSGVG	SLTMAAGG
↔ AAV8								
↔ AAV9 ↔	· · · ·							
↔ 300	230 310	240 320	250 330	260	270	280	290 _	
AAV-DJ ⇔VGNSSGNWHCE AAV2	STWMGDRVITTS	TRTWALPTYNN	NHLYKQISNSTS	GGSSNDNAYF	GYSTPWGYFD	FNRFHCHFSPI	RDWQRLINNNWGFR	PKRLSFKI
AAV8 ↔	•••							
AAV9 ↔								
↔ 410	340 420	350 430	360 440	370	380	390	400 _	
AAV-DJ →TKTIANNLTSI AAV2 AAV8	LIQVFTDSEYQLP	YVLGSAHQGCI	LPPFPADVFMIP	QYGYLTLNNG	SQAVGRSSFY	CLEYFPSQMLI	RTGNNFQFTYTFED	VPFHSSYA
·····································								
↔ 520	450 530	460 540	470 550	480	490	500	510 _	
AAV-DJ →DQYLYYLSRTÇ AAV2	2 TTGGTTNTQTLG	FSQGGPNTMAN	NQAKNWLPGPCY	RQQRVSKTSA	DNNNSEYSWT	GATKYHLNGRI	DSLVNPGPAMASHK	DDEEKFFP
⇔AAV8		A						
↔ 630 AAV-DJ	560 640	570 650	580 660	590	600	610	620 🔒	
	TDEEEIRTTNPV	ATEQYGSVST	NLQRGNRQAATA	DVNTQGVLPG	MVWQDRDVYL	QGPIWAKIPH:	FDGHFHPSPLMGGF	GLKHPPPQ
↔ AAV8 ↔							_	
	670	680	690	700	710	720	730	
AAV-DJ →TTFNQSKLNSF AAV8	TITQYSTGQVSVE	IEWELQKENS	KRWNPEIQYTSN	YYKSTSVDFA	VNTEGVYSEP	RPIGTRYLTRI	JL	
\hookrightarrow								

Same map from the corresponding -par.html file, using default colors:



5.3 Sequence definitions

The -def.txt file shows comprehensive definitions of variant sequences in terms of their parent(s), using as few instructions as possible. This file is most useful for designing synthesis, PCR and assembly strategies for the reconstruction of the variants from the parents. For each variant, a 5-column table is displayed:

- Variant region: Each region is defined as a range (ex. 86-147), a number (ex. 457) or two consecutive numbers separated by a slash (ex. 233/234). In the first case, the numbers represent sequence positions, and both are included in the region (ex. the 1-10 region includes all 10 first positions in the sequence). In the second case, the region is a single position. It the third case, the region is the empty interval between two consecutive positions, and shows the presence of a deletion.
- 2. **Parent/feature:** In the case of sequence identity with a parent, the parent name is displayed. Otherwise, the name of the feature (substitution, insertion, deletion) is displayed. Unmatched regions that can not be assigned to a parent are shown as "unmatched".
- 3. **Parent region:** The corresponding region in the parent sequence. For substitution, insertions and deletions, the parent is the one surrounding the feature (or next to the feature if the feature is at a sequence end). Position numbering is the one defined in the parental sequence file as explained in parents (first position in the sequence becomes the number next to the sequence name if a number if present).
- 4. **Variant sequence:** When a substitution, insertion or unmatched region is present, the region sequence is displayed here.
- 5. **Parent sequence:** In the case of substitutions or deletions, the corresponding parental sequence is displayed here.

Here is an example of a -def.txt file:

```
Variant name: OLIG001
Variant region Parent/feature Parent region Variant sequence Parent sequence``
1-124 AAV2 1-124
```

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F				
86-156	AAV9	86-156		
136-178	AAV6	136-178		
165-262	AAV9	165-262		
234-341	AAV8	235-342		
333-373	AAV2	331-371		
343-531	AAV8	344-532		
532	substitution	533	K	E
533-725	AAV8	534-726		
726	unmatched		Н	
727-737	AAV6	726-736		
Variant name: A	AV2i8			
Variant region	Parent/feature	Parent region	Variant sequence	Parent sequence
1-584	AAV2	1-584		
582-590	AAV8	585-593		
591-735	AAV2	591-735		
Variant name: r	AAV2-retro			
Variant region	Parent/feature	Parent region	Variant sequence	Parent sequence
1-381	AAV2	1-381		
382	substitution	382	D	N
383-587	AAV2			
588-597	insertion	587/588	LADQDYTKTA	
598-717	AAV2	588-707		
	substitution		-	V
719-745	AAV2	709-735`	×	

5.4 Alignments

The -aln.txt file consists of two parts: the first concerns sequences derived from a single parent, while the second part is about sequences derived from multiple parents. In each part, variants derived from the same main parents are aligned together against their main parent. Only regions that differ from the parent in any of the variants are shown, and the region position is indicated by a number (numbering is the one defined in the parental sequence file as explained in parents). Sequence identity is shown using the identity symbol (default: '.'), gaps are shown using the gap symbol (default: '-'). Insertions are indicated by a gap in the parent sequence, deletions by a gap in the variant sequence. For substitutions, the actual sequence is shown. Increasing the region area (using the -e argument) allows to show more sequence context and to combine neighboring regions. The -aln.txt file is most useful for variants derived from a single parent (such as from insertion or targeted substitution libraries), as it allows to compare variants in discrete regions, which can be much larger in number and size when variants are derived from multiple parents (as in shuffled libraries). Example of -aln.txt file:

```
Alignment of variant sequences against their main parental sequence - single parent
             239 251 262 271 381 443 449
                                            458
                                                  491
                                                        498
                                                                506 587
\leftrightarrow 699 703 707 729
             VIT TYN SQS HYF NNG YYL NTPSGTT SRLQF TSADN SEYSWTG TKY N-----R.
AAV2
→QYT NYN SVN RYL
LI-A
             ... .A. ... .F. .SEGSL. .S.G. .DGE. .DF.... ... .-----.
\hookrightarrow . . . . . . . . . .
             LI-C
\hookrightarrow . . . . . . . .
AAV2-sept-Y-F ... .F. ... .F. ... .F. .... .F. .... .F. ... .F. ... .F. ... .F. ... .F. ... .F.
\hookrightarrow .F. ... .F.
```

(continues on next page)

7m8 .T.LALGETTRPA. ... ↔I. ... rAAV2-retroD.D.LADQDYTKTA. ... \hookrightarrow I. ... 588 Q----A AAV9 AAV-F .FVVGQSY. AAV-PHP.B .TLAVPFK. Alignment of variant sequences against their main parental sequence - multiple parents 124 134 161 167 178 187 197 204 232 261 270 309 326 → 339 371 409 448 455 AAV2 RVL EPVK KAG ARK DADS PLGQP GLGTNT ATGSG STWMG S-SQS-GAS NHY KRLNF. ↔NDGTTT TVQ MVP FTFSY TNTPSG TTQSRLQF AAV2i8 AAV-DJ .L. .AA.I.E. .V.SL. .A.G.N.T.G.S. .A. ...S. .E.. \hookrightarrow K. .I. .I. .Q.T. .Q.TG. .NTQT.G. NP84 531 584 655 661 464 704 712 AAV2 QAGASDIRDQSRN EKF QRGNRQAA ANPST FSAAKFAS YNKSVNV FTVDTNG AAV2i8QQ.TAP. AAV-DJE. .G...... NP84 29 33 49 54 59 65 1 12 75 ↔ 101 114 580 MSFVDHPP E-EVGEGLREFLGLEAG PKP PNQQHQDQAR YNY PGN DRG NRADEVARE. AAV5 →DISYNEQLEAG EKLADD GKA PAT AAV2.5T .AADGYL. .DTLS..I.QWWK.KP. .P. .AER.K.DS. .K. .F. .K. .E..AA.L. .KA.DR.. →DS. .R.KE. .R. .T. 13 20 66 80 91 104 124 134 147 151 157 2.8 →162 168 179 188 200 205 211 233 342 532 726 AAV8 DNL REWWAL GAPKPKANQQKQDDGR NAA DQQLQAG LRY LQE RVL EGA EPS QRSP STG_ ↔KKG ARK DSE PLG GPNT AAG PMA STW TIQ DEE PRP OLIG001 .T. .Q..K. .P.P..PAERHK..S. .E. .R..DS. .K. .K. .L. .A. .Q. .-E. .S. .T. → .K. .T. .I. .SL. .S. .V. .Q. .V. .K. .H. 736 AAV8 RNL OLIG001 .Р. 23 30 37 55 91 104 124 313 326 341 415 448 ш. 482 490 _ WAL POP OHODNAR PGN LKY LKE RLL LNF TDNNGVK TVO YEFENV SKTINGSG-AAV9

→QNQQTLKFSVAGPSNMAVQGRNYIP PSY STTVTQN

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SCH2 .D. .K. .K..DG. .F. .R. .Q. .V. .S. .Q.E.T. .I. .T..D. .R.NTP..TTT.SR.Q. \hookrightarrow .Q..A.DIRD.S..WL. .C. .K.SAD. SCH9 $\hookrightarrow \ldots \ldots \ldots \ldots . \mathbb{WL} \ldots \mathbb{C} \ldots \mathbb{K} . \mathbb{SAD} .$ 500 514 528 536 547 554 561 566 575 AAV9 EFAWPGASSWAL RNSLMN KEGEDRF PLSGSL GTGRDN DADK TNE IKT →ESYGQVATNHQSAQAQAQTGWVQNQGIL SCH2 .YS.T..TKYH. .D..V. .DD.EK. .Q..V. .SEKT. .IE. .D. .R. .Q..S.S..L.RGNR.. →A.AD.NT..V. SCH9 .YS.T..TKYH. .D..V. .DD.EK. .Q..V. .SEKT. .IE. .D. .R. .Q..S.S..L.RGNR.. →A.AD.NT..V.

Examples

In order to explore the functionalities of parent-map, a set of protein sequences of evolved or designed AAV capsid variants described in peer-reviewed articles was assembled, as well as a set of the protein sequences of all wild-type parental AAV capsids from which they are derived. They can be downloaded by clicking on the following links:

Variants

Parents

Variants SCH2 and SCH9 are derived from a chimeric library combining fragments from AAV2, AAV4, AAV5, AAV6, AAV8 and AAV9. Using default values, parent-map correctly identifies parental contributions from AAV2, AAV6, AAV8 and AAV9 for both variants.

Variant AAV-F derives from an AAV9 peptide insertion library. Parent-map correctly identifies AAV9 as the single parent and detects peptide sequence FVVGQSY insertion between positions 588 and 589.

Variant AAV-DJ derives from a shuffled DNA library made with parental serotypes AAV2, AAV4, AAV5, AAV8 and AAV9. Parent-map correctly identifies parental contributions from AAV2, AAV8 and AAV9, and detects an A to T substitution at position 457 (corresponding to position 458 in parent AAV8).

Variant AAV-PHP.B derives from an AAV9 peptide insertion library. Parent-map correctly identifies AAV9 as the single parent and detects peptide sequence TLAVPFK insertion between positions 588 and 589.

Variant AAV2.5T derives from a capsid library generated by DNA shuffling of parents AAV2 and AAV5 and errorprone PCR. Parent-map correctly identifies parental contributions from AAV2 (VP1-specific N-terminal region) and AAV5 (remainder of the sequence) and an A to T substitution at position 582 (581 in parent AAV5).

Variant 7m8 derives from an AAV2 peptide insertion library. Parent-map correctly identifies AAV2 as the single parent and detects peptide sequence LALGETTRPA insertion between positions 587 and 588 (AAV2 numbering), as well as substitutions I to T at position 240 and V to I at position 718 (708 in AAV2).

Variants LI-A and LI-C derive from a synthetic library based on AAV2 with mutations at targeted positions. Parentmap correctly identifies AAV2 as the single parent, as well as the 14 and 4 substitutions in LI-A and LI-C respectively.

Variant NP84 derives from a shuffled DNA library. Parent-map describes it as combining regions of AAV2, AAV3B and AAV6, as well as having substitutions K to E at 532 and R to G at 585 in the C-terminal AAV2-homolog region.

Variant OLIG001 derives from a shuffled DNA and error-prone PCR library. Parent-map identifies parental contributions from AAV2, AAV6, AV8 and AAV9, as well as a E to K substitution at position 532 (position 533 in relevant parent AAV8) and an unmatched H at position 726 between regions homologous to AAV8 and AAV6.

Variant AAV2i8 is a rationally designed chimeric capsid based on AAV2 in which the receptor footprint region was replaced with corresponding region from AAV8. Parent-map successfully identifies AAV2 and AAV8 parental contributions.

Variant rAAV2-retro derives from an AAV2 peptide insertion and error-prone PCR library. Parent-map correctly identifies AAV2 as the parent, the peptide sequence LADQDYTKTA insertion between positions 587 and 588, as well as the 2 substitutions N to D at position 382 and V to I at position 718 (708 in AAV2).

Variant AAV2-sept-Y-F is a rationally designed multiple mutant of AAV2 combining seven Y to F mutations on its surface. Parent-map correctly identifies AAV2 as the single parent, as well as the 7 Y to F substitutions at positions 252, 272, 444, 500, 700, 704 and 730.

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