The T_EX**shade** package*

Typesetting nucleotide and peptide alignments

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v1.22; 2010/10/11

Abstract

Setting alignments of nucleotides and peptides for publication or presentation purposes is usually a time consuming two-step process. First, a scientific software is used for the calculation of the alignment. This is done in a few minutes. Then, in order to highlight special sequence relationships and to label positions and regions of interest a second software with high quality output capability is needed. Manipulating sequence alignments with standard word processing or graphics programs takes its time—often several hours—and simple layout changes such as re-breaking lines, say from 50 to 40 residues per line, elongate the working time considerably.

 T_EX shade is an alignment shading software written in T_EX/IAT_EX which can process multiple sequence alignments in the MSF, ALN and FASTA file format. It provides in addition to common shading algorithms special shading modes featuring functional aspects, e.g. charge or hydropathy, and a plenitude of commands for handling shading colors, text styles, labels, legends and even allows the user to define completely new shading modes. T_EX shade combines highest flexibility and the habitual T_EX output quality—with reasonable time expenditure.

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Contents

1	Pac	kage Overview 4	2
	1.1	Version History	2
	1.2)
		1.2.1 Typesetting documents using)
		1.2.2 Memory shortness when using T _F Xshade 12	2
	1.3	System requirements	c
	1.4	The texshade environment 15)
	1.5	Shading modes predefined in this package)
		1.5.1 Identity mode	,
		1.5.2 Similarity mode	;
		1.5.3 T-Coffee shading 19)
		1.5.4 Diversity mode \ldots 19)
		1.5.5 Functionality modes)
	1.6	Bar graphs and color scales	,
	1.7	Secondary structures)
	1.8	Sequence fingerprints	
	1.9	Sequence logos	
	1.10	Subfamily logos	
		Customization of the alignment output	
2	For	mat of alignment input files 38	;
	2.1	The MSF file format 38	;
	2.2	The ALN file format)
	2.3	The FASTA file format	1
3	Use	of a T _E Xshade parameter file 43	;
4	texs	shade user commands 44	L
	4.1	Using predefined shading modes	E
	4.2	Creating new functional shading modes	
	4.3	Appearance of the consensus line	
	1.0	4.3.1 Residue weight tables	
	4.4	Display of logos	
	1.1	4.4.1 Sequence logos	
		4.4.2 Subfamily logos	
	4.5	Appearance of the sequence lines	
	1.0	4.5.1 Names, numbers and gaps	
		4.5.2 Displaying selected residues in the alignment	
		4.5.3 Hiding, killing, separating and ordering 61	
		- i.o.o - inding, mining, separating and ordering	

		4.5.4	Residues per line and further settings	6	52
		4.5.5	Fingerprinting	6	53
	4.6	Individ	lual shading and labeling of sequence stretches .	6	53
		4.6.1	Shading of regions and blocks	6	64
		4.6.2	Emphasizing, tinting, lowercasing, and framing.	6	66
		4.6.3	Graphical labeling of sequence features	6	67
		4.6.4	Including secondary structure information	7	75
	4.7	Displa	ying and building legends	7	78
	4.8	Adding	g captions to the alignment $\ldots \ldots \ldots \ldots \ldots$	7	79
	4.9	Font h	and ling \ldots \ldots \ldots \ldots \ldots \ldots \ldots \ldots	7	79
		4.9.1	Changing font styles	7	79
		4.9.2	0 1		31
	4.10	Goodie	es	8	31
		4.10.1	Molweight and charge	8	32
		4.10.2	Similarity/identity data and tables	8	32
5	The	PostS	cript color selection scheme	8	34
6	\mathbf{List}	ing of	the texshade default settings	8	87
	6.1	Standa	rd definitions	8	37
	6.2	Colors	used in the different shading modes	8	37
	6.3	Residu	e weight tables	Q	91
7	Qui	ck Ref	erence	9	6
8	Refe	erences	3	10)4

1 Package Overview

After texshade.ins is run through T_EX the following files should appear in the directory:

texshade.sty	the style file with all T _E X <mark>sh</mark> a d e commands
texshade.def	an example parameter file with the standard
	parameter settings
AQPDNA.MSF	an example nucleotide alignment (MSF-format)
AQPpro.MSF	an example protein alignment (MSF-format)
AQP_TC.asc	an example T-Coffee shading file (score_ascii-format)
AQP2spec.ALN	a further protein alignment (minimal ALN-file)
AQP1.phd	secondary structure information (PHD-format)
AQP1.top	topology data extracted from AQP1.phd
AQP1_HMM.sgl	topology information (single line, HMMTOP-format)
AQP1_HMM.ext	topology information (extended, HMMTOP-format)
standard.cod	standard genetic code definitions
ciliate.cod	ciliate macronuclear genetic code

The alignment file examples as well as the topology data file are needed for T_EX ing this documentation and can serve as illustrations for the MSF and ALN file format.

The following subsections give an overview on the capabilities of the $T_E X$ shade package. All commands are described in detail later on.

1.1 Version History

v1.22 2010/10/11

Introductions: Sets of shading colors can be defined and named using \defshadingcolors (p. 45) and different sets of colors can be selected for different sequence blocks by \changeshadingcolors (4.6.1).

v1.21 2010/03/01

Corrections: \setdomain now recognizes a changed \startnumber.

Introductions: (a) Sequence motifs (4.6.1) can be given as a selection pattern in sequence stretch labeling commands, such as \shaderegion, \emphregion or \frameblock, instead of numeral stretch definitions.

(b) A new type of sequence emphasis by lowercase letters was implemented: \lowerregion and \lowerblock (4.6.2).

v1.20a 2009/11/30

Corrections: Identity/similarity tables work now without having to set a conservation plot in the feature line.

v1.20 2009/10/05

Corrections: Landscape mode works now correctly. Annoying **Overfull \hbox** ... signals appearing after **\residuesperline*** commands are ignored; domains work now with frequency corrected sequence logos.

Introductions: (a) New commands for setting residue weight tables were introduced \weighttable (with parameters: identity, structural, PAM250, PAM100, and BLOSUM62), \gappenalty, and \setweight. (b) Similarity/identity percentage tables can be printed with \similaritytable¹ and values for specific sequence pairs can be utilized with \percentsimilarity{ $\langle seqref2 \rangle$ } and \percentidentity{ $\langle seqref2 \rangle$ }.

v1.19a 2009/06/11

Correction: Gaps are now treated correctly when setting domains.

Introduction: Stop-positions in protein sequences (* in the input file) can be indicated by a special character (\stopchar) in the alignment output.²

v1.19 2009/03/09

Correction: Logo characters are now horizontally scalable using \charstretch. Minor bugs were fixed.

Introduction: (a) Selection of residue positions was enhanced (4.6.1) T_EX **shade** can now select residues based on their 3D coordinates provided in a PDB structure file. 3D selection can be due to a certain distance around a point, along a line, or above and below a plane. It works with \feature, \shaderegion, \shadeblock, \tintregion, \tintblock, \emphregion, \emphblock, \frameblock. (b) The list of selected residues is printable with \printPDBlist or viewable during the T_EX run with \messagePDBlist. (c) \hideblock and related commands were replaced by \setdomain (4.5.2). This command will display only selected residues in the alignment. Thickness and colors of a domain separator rule can be set using \domaingaprule and \domaingapcolors.

 $^{^1\}mathrm{Asked}$ for by Giovanni M. Lesa.

²Suggestion by Yun He.

v1.18 2008/04/15

Correction: several bug fixed concerning featurename display, sequence ordering and numbering.

Introduction: (a) T-Coffee shading information can be loaded and put on the alignment.³ The conservation data can also be displayed in the consensus as well as feature color scales and bar plots. (b) Two more feature lines were added on the top and at the bottom (tttop, tttop, bbbottom, bbbbottom). (c) The startnumber and setends commands have been fused; either command can set both, a new start number as well as end definitions of the sequence section to be displayed.

v1.17 2007/06/19

Introduction:⁴ (a) A second threshold percentage was introduced in order to label two levels of conservation in 'identical' and 'similar' mode. This is achieved by setting an optional parameter in \threshold or in \allmatchspecial, or by using a number as an optional parameter in \shadingmode. (b) The feature lines can be additionally labeled with a name left or right of the feature. This is handled using \showfeaturename, \showfeaturestylename, \hidefeaturename, \hidefeaturestylename, \hidefeaturenames, \hidefeaturestylenames. The color of such names can be featurenamecolor,changed with \featurestylenamecolor, \featurenamescolor. \featurestylenamescolor, Font styles in feature names be set can as usual, e.g. \setsize{featurenames}{large} \featurestylenamesrm. or

v1.16 2007/02/18

Correction: T_EX shade crashed when calculating conservation using sequences with untypical residue characters, such as "X". Fixed. The reference sequence in diverse mode can now be shaded with \conservedresidues and, if active, \allmatchresidues .⁵

Introduction:⁶ (a) A command was introduced, i.e. $\ensuremath{\backslash} exportconsensus$ which produces a pymol script file for coloring a 3D model according to $T_EXshade$'s conservation calculation.

³Suggestion by Florian Mertes.

⁴Asked for by Marat Kazanov.

⁵For this and suggesting namerulerpos credit to Marco Pasi.

⁶Both extensions were suggested by Phillip Hahn.

(b) With namerulerpos labels of the ruler can be exchanged by a string. (c) Various parts of the alignment can now be hidden by hideblock.

New home: T_EX**shad**e, T_EXtopo, and BIOT_EX have a new home: www.pharmazie.uni-kiel.de/chem/Prof_Beitz/biotex.html.

v1.15 2006/06/27

Correction: Sequence and subfamily logos can now be plotted with pdflatex; pstricks is not needed anymore.

v1.14 2006/05/11

Introduction: In order to better recognize relevant positions in a subfamily logo [14], a bit-value can now be set by **\relevance** above which a deviation is considered relevant. Such positions can be labeled with a symbol by **\showrelevance** and hidden by **\hiderelevance**.

v1.13 2006/02/23

Corrections: Helix symbols in feature lines were not drawn correctly if the standard Computer Modern Font was changed to another one, e.g. Palatino.⁷ Fixed. Unintended gaps occurred due to numbers at the end of lines in Clustal W alignment files. Fixed. The limitations in the number of sequences per alignment have finally been overcome by a more restrictive use of counter variables.

Introductions: (a) The numbering can now be displayed—in addition to left or right—on both sides of the alignment with the optional parameter {leftright} in the \shownumbering command (p.56). (b) TeXshade tries to guess the sequence type, i. e. protein or nucleotide, if not defined by the user. (c) Plotting of sequence logos has been implemented (p.54). Logos can be shown in addition to or together with the consensus, or alone without any alignment sequences. (d) The ruler numbering can be rotated in order to make labeling of every position possible. (e) A new way to visualize subfamily characteristics has been implemented, i.e. subfamily logos (p.56) [14].

v1.12 2005/09/20

Corrections: When regional labeling with \shaderegion, \emphregion, \tintregion, or \frameblock was combined with

⁷Thanks to Markus Heller

\setends incorrect output was produced lacking the labeling.⁸ Other minor fixes.

Introductions: An additional optional parameter for setting consensus colors was implemented in the \showconsensus command (p.51). This even allows one to use color scales illustrating sequence conservation in the consensus line.

v1.11 2005/04/13

Corrections: Bounding boxes with \frameblock had a wrong height when \separationlines were used. Other minor fixes.

Introductions: (a) An additional parameter for setting individual bar and arrow thicknesses in feature lines has been introduced. (b) Additional parameters for setting the frame color and thickness of boxes in feature lines have been implemented. (c) Three more color scales have been defined: RedBlue, RedGreen, and HotCold. (d) Plotting of amino acid features (hydrophobicity, molweight, charge) as bar graphs or color scales. (e) Plotting of protein sequence conservation as bar graph or color scale⁹. (f) Color scales can be used for shading the consensus sequence according to protein sequence conservation. (g) Separate command for stretching color scales \colorscalestretch.

v1.10 2005/03/29

Corrections: Plotting of color scales and bar graphs has been sped up by more than a factor of $10.^{10}\,$

Introductions: (a) More colors have been introduced, i.e. even lighter versions of the existing PostScript colors 'LightLight' plus color name and 'LightLightLight' plus color name. (b) Sequence stretches and blocks can be tinted for labeling purposes \tintreqion, \tintblock and \tintdefault. (c) A new feature label style {restriction} has been introduced. (d) Java-typical 'NaN' values are now allowed in data files for bar graphs and color scales.

v1.9 2005/02/08

Corrections: T_EX shade version 1.8 introduced an incompatibility with T_EX topo. This problem was identified by Meike Schmedt and has been fixed.

⁸Discovered by Chris Page.

⁹Ahmad Mirza asked for (e) and (f), great suggestion!

¹⁰This and (d) I owe again to Christoph Gille.

Introductions: (a) A short version of the figure caption can now be defined for display in the list of figures¹¹ \shortcaption{ $\langle text \rangle$ }. (b) A colored frame can be drawn around a sequence block for labeling purposes with the command \frameblock.¹² (c) A new look for feature arrows has been implemented with scalable line thickness and a new end style 'ball'. (d) HMMTOP topology predictions can now be included for plotting feature lines with information on the location of the transmembrane domains.¹³

v1.8 2004/08/26

Corrections: Only minor bugs were fixed.

Introductions: (a) More colors have been designed, i.e. 'light' versions of the existing PostScript colors. (b) Three color ramps in 5% steps have been introduced: i) Blue-Red, ii) Green-Red and iii) Cold-Hot. (c) Two new feature label styles **bar** and **color** have been introduced which allow one to display number values as bar graphs or color scales along the alignment¹⁴.

v1.7 2004/01/05

Corrections: Several bugs were fixed. In gaps the wrong character was plotted in 'donotshade' mode. Gaps were colored incorrectly when a single sequence was set as consensus. Another 'donotshade' problem was solved which led to a halt of the LaTeX run¹⁵. Due to several requests, the gap and match labels in **diverse** mode were switched ('-' in gaps; '.' at matching positions) in order to follow convention. Introduction: T_EX**shade** speaks spanish (\spanishlanguage). Necessary translations were contributed by Mikel Egaña Aranguren. A new feature label style helix has been introduced.

v1.6 2002/03/26

Corrections: The unnecessary restriction to the DVIPS driver for color.sty has been removed¹⁶. Any color.sty compatible driver option can be given with the \usepackage{texshade} call and is then passed to the color package. The '\namecolor' and '\numbercolor' commands do now support sequence lists.¹⁷

¹¹ Meike, here you go ...

 $^{^{12}\}mathrm{Alan}$ Robinson, this is for you.

 $^{^{13}\}mathrm{Implemented}$ after a request by Steffen Moeller.

¹⁴Inspired by Christoph Gille's STRAP

 $^{^{15}\}mathrm{Thanks}$ to Jeferson J. A renzon and Naomi Siew

¹⁶As suggested by Eckhart Guthöhrlein.

¹⁷Thanks to Denys Bashtovyy.

Introductions: (a) The FASTA file format is supported by T_EX shade as alignment inputs. (b) Two commands set the space between sequence blocks either to be flexible (as so far) '\flexblockspace' or the be fixed '\fixblockspace'. (c) One can now refer to sequences by their name in addition to the number in the input file. (d) Using '\firstcolumnDSSP' and '\secondcolumnDSSP' one can choose which of the first to columns should refer to the sequence numbering (the second column remains default setting)¹⁸.

v1.5a 2001/03/08

Corrections: 'X's in the alignment file caused a run-time error. Fixed. Introductions: (a) The vertical space between feature lines can be controlled by four new commands: \ttopspace, \topspace, \bottomspace and \bbottomspace¹⁹. (b) It is now easily possible to add a caption to the alignment with the \showcaption command. (c) T_EXshade stores the sequence lengths in the .aux file in order to have correct breaks of the gaps after the sequences.

v1.4&4a 2000/9/12 & 2000/10/3

Introductions: (a) The alignment legend can now be moved by the command '\movelegend'. (b) In commands with parameters that contain series of sequence numbers, e.g. \orderseqs, a dash can be used, e.g. {1-3,6-4,7} instead of {1,2,3,6,5,4,7}.

v1.3a&b 2000/7/28 & 2000/7/30

Introductions: (a) It is now possible to force T_EX shade to display gap symbols before and after the actual sequence by the commands '\showleadinggaps' and '\hideleadinggaps' (4.5.1). (b) The sequence names input routine is now more tolerant concerning special characters.

v1.3 2000/3/3

Corrections: Line scrambling occured when features where set in the ttop row without a feature in the top row. Fixed. The incompatible command '\language' with the babel package has been replaced by '\germanlanguage' and '\englishlanguage'²⁰.

¹⁸c and d were suggested by Christoph Gille.

¹⁹Suggested by Ulrike Folkers.

²⁰Thanks to Eckhart Guthöhrlein.

Introductions: (a) Now, translations of sequence stretches are possible. Either nucleotide or amino acid sources can be translated. This is done by the new {translate} option for the feature command. (b) The codons are defined by the new command '\codon'. Complete codon sets can be loaded by '\geneticcode'. (c) Further, the size and style of the nucleotide triplets of backtranslations can be set by '\backtranslabel' and '\backtranstext'. (d) Two more feature counter styles were introduced: '\Romancount' and '\romancount'. (e) TEXshade is now compatible with TEXtopo, a new TEX software for drawing and shading topology plots of membrane proteins.

$v1.2a \ 1999/6/24$ (not released)

Minor corrections: '\namecolor' and '\numbercolor' are now really correctly reordered. Brackets (and) are now allowed in sequence names. The option {case} in '\funcshadingstyle' works now.

v1.2 1999/6/12

Corrections: (a) Functional group definitions of more than seven groups produced an error when displaying group number eight. These residues where skipped in the alignment. Fixed.

Introductions: (a) Protein secondary structure files in the DSSP, STRIDE and PHD format can be included and displayed automatically within the alignment by '\includeDSSP' (and similar commands for STRIDE, PHDsec and PHDtopo, 4.6.4). (b) Which types of secondary structures are to be included or skipped in the alignment is chosen by '\showonDSSP' and '\hideonDSSP' (and respective commands for STRIDE, PHDsec and PHDtopo). (c) The appearance of the labels is defined by '\appearance'. (d) Internal counters for repeatedly occuring structure types can be activated by '\numcount', '\alphacount' and '\Alphacount'. All commands are described in 4.6.4.

v1.1 1999/5/26

Corrections: (a) The activation of 'emphregion' lead to an emphasized following alignment. This has been corrected. (b) '\namecolor' and '\numbercolor' were not reordered with the command 'orderseqs'. Fixed. (c) Sequence gaps at the beginning or the end of a sequence, i.e. before the first and after the last residue where labeled with the gap symbol. Now these positions are left blank.

Introductions: (a) In order to treat the preceeding and sequence following gaps correctly, T_{FX} shade needs to know the length of the sequences. Therefore, the command '\seqlength' was introduced (4.5). (b) With '\gapcolors' (also 4.5) the color selection for gap symbols is independent from non conserved residues. (c) The divisions of the ruler where so far fixed to 10. Now, this value is changeable by '\rulersteps' (again 4.5). (d) '\hideresidues' and '\showresidues' turn off or on the residue names, i.e. one can choose between a display of shaded boxes only or with letters in the boxes (4.5.3). (e) The changes (c) through (d) were necessary for the introduction of '\fingerprint'. This command allows one to display the complete sequence in one line for an easy survey of the alignment (4.5.5).

v1.0 1999/5/12

First release.

1.2 I_{E} basics

1.2.1 Typesetting documents using PT_EX

In order to use any of the macros provided by the BIOTEX-project (TEXshade/TEXtopo) efficiently a basic understanding of the TEX typesetting system and its usage is required. Several books are available on this topic, but a rather quick and easy introduction is the Not so short introduction to ETEX. This document is available from all Comprehensive TEX Archive Network (CTAN) servers, e.g. from ftp://ftp.dante.de/pub/tex/documentation/lshort/, in many different languages and formats besides ETEX, such as POSTSCRIPT and on-line viewable PDF. I also put a link from the BIOTEX (TEXshade/TEXtopo) homepage to the document collection (http://pharmazie.uni-kiel.de/chem/Prof_Beitz/BioTeX).

1.2.2 Memory shortness when using T_EXshade

If you are using T_EX**shade** to align several large sequences (about 1000 residues/sequence), LaTeX will probably stop compiling and quit with one of the following messages:

!\ TeX capacity exceeded, sorry [main memory size=384000] or

!\ TeX capacity exceeded, sorry [stack size=300].

 $T_{E}X$ allocates space for different kinds of internal variables. Setting alignments needs lots of memory, usually more than for typesetting

plain text. Thus, the parameter settings of a standard T_EX installation might not be sufficient for certain projects. This manifests in T_EX error messages about insufficient memory and the setting process is interrupted. There is no reason to be concerned. The parameters can be set by hand. Unfortunately, each T_EX system hides its default parameter file in a different place in the system.

In the following, an excerpt from the FAQ-list to T_EX**shade** is added. This explains how to increase the settings in OzT_EX for the Macintosh, MikT_EX for Windows and teT_EX for *NIX T_EX distributions. Please contribute to this list!

1. OzT_EX 4.0 for the Macintosh:

Find the file 'OzTeX:TeX:Configs:Default'. This file contains all memory settings. Look for the section '% TeX parameters' and increase the values that T_EX complains about during the run. You will have to restart OzT_FX before the changes are active.

For older versions of OzT_EX the configuration file has the same name but the path is somewhat different.

2. teT_EX for *NIX: (contributed by Joerg Daehn)

Find the file: '/usr/share/texmf/web2c/texmf.cnf' or use

locate texmf.cnf at the command prompt to find it.

Login as super user. Backup 'texmf.cnf' in case you destroy something and then open the 'texmf.cnf' file in your favorite text editor and use its search function to locate main_memory. This variable is set to 384000. Change this to some higher value, i.e. 4000000 (works fine for me!). The total amount of memory should not exceed 8000000, so check the other values in that section.

Next, you want to change the stack size. Search for stack_size. This will be set to 300. I changed it to 4000 and it works fine.

There might be complains by T_EX about further specific parameters such as stack_size. You find all those in the same file.

After this you have to run 'texconfig init'.

Logout as root.

After this all should be set for large alignments. Happy T_EXing!

The information on how to achieve this was derived from a mail in the teT_EX mail archive. The original question was posted by Pascal Francq and answered by Rolf Nieprasch.

3. MiKT_EX for Windows:

The $MiKT_EX$ documentation describes very detailed how the memory settings can be changed. In brief, you must locate the configuration file 'miktex/config/miktex.ini'. In the [MiKTeX] section of this file you find all the parameters you need, e.g. mem_min, mem_max, buf_size, stack_size etc.

It appears, that the standard settings of $MiKT_EX$ are bigger than that of other T_EX installations, so it may not always be necessary to increase the values.

1.3 System requirements

TEX shade requires $\[mathbb{L}^{*}T_{E}X 2_{\varepsilon}\]$ with color.sty and graphics.sty for shading. For arrows in the feature line (p.67) the AMS Math style is needed. David Carlisle's color.sty is part of the Standard $\[mathbb{L}^{*}T_{E}X\]$ 'Graphics Bundle' [1]. This and the other packages can be downloaded from any TEX archive, e.g. ftp.dante.de; usually they are included in a comprehensive TEX installation.

The color style allows one to use several $[\langle options \rangle]$, e.g. dvips, pdftex or dviwin. These provide the commands which different devices/programs need to display colored output. It is advisable to make yourself familiar with the color.sty manual. You should define a default driver in the file color.cfg. Since there is no direct call of color.sty by the user, the option can be stated when T_EXshade is loaded, see next subsection. If no option is stated the DVIPS driver will be loaded.

With the [dvips] option the output DVI-file can be converted to POSTSCRIPT using the DVIPS program and can later be viewed or printed with the public domain GHOSTVIEW program which is available for almost all computer platforms. Further, more and more standard T_EX viewers are to a certain extent POSTSCRIPT compatible.

1.4 The texshade environment

The commands provided by the T_EX_{shade} package are enabled by the following command in the document header section:

Make sure that the file 'texshade.sty' is present in a directory searched by T_EX (see the installation notes in the file 'texshade.txt'). The $\langle option \rangle$ given here is passed to color.sty which handles the color commands for a particular output device, see previous subsection and the color.sty manual.

The T_EX**shade** package provides only one single new environment: texshade. This environment has one mandatory and one optional argument, both of them designating file names which must be present in a directory searched by T_EX. The required file $\langle alignmentfile \rangle$ contains the aligned nucleotide or peptide sequences (see section 2). This file is needed, because T_EX**shade** does no alignment by itself, it has to take a preprocessed alignment as input. The optional file is a parameter file (section 3) with definitions for the customized calculation of the consensus, special sequence features or labels etc. In this parameter file all T_EX**shade** commands which are allowed in the texshade environment can be used and are fully functional. Within the environment further T_EX**shade** commands can be given to replace or complete settings from the parameter file.

Thus, setting an alignment with $T_{E}X$ shade is as simple as this:

 $\begin{texshade}[\langle parameterfile \rangle] {\langle alignmentfile \rangle} \\further T_EX shade commands, if needed \\\end{texshade} \$

1.5 Shading modes predefined in this package

1.5.1 Identity mode

This basic type of shading is provided by almost any alignment program. All identical residues at a position are shaded if the number of matching residues is higher than a given threshold (default is 50%).

AQP1.PRO	TLGLLLSCQISILRAVMYIIAQCVGAIVASAIL	112
AQP2.PRO	TVACLVGCHVSFLRAAFYVAAQLLGAVAGAAIL	104
AQP3.PRO	TFAMCFLAREPWIKLPIYTLAQTLGAFLGAGIV	112
AQP4.PRO	TVAMVCTRKISIAKSVFYITAQCLGAIIGAGIL	133
AQP5.PRO	TLALLIGNQISLLRAVFYVAAQLVGAIAGAGIL	105

```
\begin{texshade}{AQPpro.MSF}
   \setends{1}{80..112}
   \hideconsensus
\end{texshade}
```

Quite uncommon for an alignment shading program is the possibility to display only selected sequence domains, e.g. to eliminate uninteresting positions from the output:

			10 120 130
AQP1.PRO	<u>Ť</u> LGL <mark>L</mark> LSCQ I S	AQCVGAIVASA	Á LENS <mark>LG</mark> R <mark>N</mark> DLÁ
AQP2.PRO	TVACLVGCHVS	AQLLGAVAGA	A IR <mark>GDLAVN</mark> ALH
AQP3.PRO			VS <mark>G</mark> PN <mark>G</mark> TAGIF
AQP4.PRO			VV <mark>G</mark> GLGVTTVH
AQP5.PRO	TLALLIGNQIS	AQLVGAIAGAC	ARGNLAVNALN

Code:

```
\begin{texshade}{AQPpro.MSF}
   \setdomain{1}{80..90,100..110,120..130}
   \showruler{1}{top}
   \hidenumbering
   \hideconsensus
\end{texshade}
```

This goes even furher. You can have T_EX shade select positions based on the 3D coordinates provided by a PDB file, e.g. show all residues that are within an 8 Å radius around the α -carbon of the residue at position 81:

		.103	.148	$151 \\ 152 \\ 152 \\ 153 \\ 0$. 155 . 156	$193 \\ 194 \\ 195 $. 218 . 219	. 222
AQP1.PRO	S <mark>NPAVT</mark> LGL <mark>L</mark> LS	V	Q	LCV	ΑT	PAR	IG	L
AQP2.PRO	INPAVTVACLVG	L	Q	LCI	AS	PAR	VG	Ι
AQP3.PRO	LNPAVT <mark>FA</mark> MCFL	L	Α	VCV	ΑI	PAR	VS	L
AQP4.PRO	INPAVT <mark>V</mark> AMVCT	L	Q	FΤΙ	AS	PAR	ΙG	L
AQP5.PRO	INPAITLALLIG	V	Q	LCI	SS	PAR	VG	L

```
\begin{texshade}{AQPpro.MSF}
   \setdomain{1}{point[8]:1J4N.pdb,81[CA]}
   \showruler{top}{1} \rulersteps{1}
   \hidenumbering
   \hideconsensus
\end{texshade}
```

If you like, positions where conservation is very high (here $\geq 80\%$) can be shaded in a special color and the consensus can be shown with or without shading according to the degree of conservation:

AQP1.PRO	TLGL <mark>L</mark> LSCQ IS I <mark>LRAV</mark> MYIIAQCVGAIVASAIL	112
AQP2.PRO	TVACLVGCHVSFLRAAFYVAAQLLGAVAGAAIL	104
AQP3.PRO	TFAMCFLAREPWIKLPIYTLAQTLGAFLGAGIV	112
AQP4.PRO	TVAMVCTRK <mark>IS</mark> IAKS <mark>VF</mark> YITAQCLGAIIGAGIL	133
AQP5.PRO	TLALLIGNQISLLRAVFYVAAQLVGAIAGAGIL	105
consensus	\overline{T} , \overline{A} , $\overline{1}$,, \overline{iS} , \overline{IravfY} , \overline{AQ} , \overline{IGAi} , \overline{GAgIL}	

Х	non co	onserved
Х	\geq 50%	conserved
Х	\geq 80%	conserved

Code:

```
\begin{texshade}{AQPpro.MSF}
    \threshold[80]{50}
    \setends{1}{80..112}
    \showconsensus[ColdHot]{bottom}
    \defconsensus{.}{lower}{upper}
    \showlegend
\end{texshade}
```

1.5.2 Similarity mode

In many cases it is expedient—mostly when comparing protein sequences—to shade also residues which are not identical but similar to the consensus sequence. Consider a position where three out of five residues are basic arginines and two more residues are also basic but lysines. In similarity mode T_EX shade shades similar residues in a different color to distinguish them from the consensus residue. Even when none of the residues alone reaches the threshold but a group of similar residues does these are shaded in the 'similarity' color. This case is given for instance when at a position in a five sequence alignment two aliphatic values and two also aliphatic isoleucins are present and the threshold is set to 50%. Neither residue exceeds this percentage but as a group of similars they do.

In grayscale printouts some colors of the following alignment may appear undistinguishable. Don't worry if you usually use grayscale all colors/grays can be selected freely (see 5).

	first case (see text)	
	\downarrow	
AQP1.PRO	TLGLLLSCQISILRAVMYIIAQCVGAIVASAIL	112
AQP2.PRO	TVACLVGCHVSFLRAAFYVAAQLLGAVAGAAIL	104
AQP3.PRO	TFAMCFLAREPW <mark>IK</mark> LPIYTLAQT <mark>LGAFLGAGIV</mark>	112
AQP4.PRO	TVAMVCTRKISIAKSVFYITAQCLGAIIGAGIL	133
AQP5.PRO	TLALLIGNQISLLRAVFY <mark>V</mark> AAQL <mark>VGAIA</mark> GAGIL	105
	second case (see text)	

Code:

```
\begin{texshade}{AQPpro.MSF}
   \shadingmode{similar}
   \threshold[80]{50}
   \setends{1}{80..112}
   \hideconsensus
   \feature{top}{1}{93..93}{fill:$\downarrow$}{first case (see text)}
   \feature{bottom}{1}{98..98}{fill:$\uparrow$}{second case (see text)}
\end{texshade}
```

Probably you know this kind of shading from the public domain program BoxShade by KAY HOFMANN or from the Macintosh version MacBoxShade by MICHAEL D. BARRON. TEXshade provides the same functionality—and goes truly beyond—for the TEX community.

1.5.3 T-Coffee shading

 T_EX **shade**'s capabilities of calculating alignment shadings are limited. T-Coffee (www.tcoffee.org) is a sophisticated alignment/shading software. You can apply shading from T-Coffee in T_EX **shade** by loading the shading information file (score_ascii) generated by T-Coffee.

feat-cons		
AQP1.PRO	GSALGFNYPLERNQTLVQDNVKVSLAFGLSIATL	63
AQP2.PRO	GSALQWASSPPSVLQIAVAFGLGIGIL	55
AQP3.PRO	GSVAQVVLSRGTHGGFLTINLAFGFAVTLA	63
AQP4.PRO	GSTINWGGSENPLPVDMVLISLCFGLSIATM	84
AQP5.PRO	GSALKWPSALPTILQISIAFGLAIGTL	56
consensus		

Code:

```
\begin{texshade}{AQPpro.MSF}
    \shadingmode[AQP_TC.asc]{T-Coffee}
    \setends{1}{30..63}
    \feature{top}{1}{30..63}{color:conservation[T-Coffee]}{}
    \showfeaturestylename{top}{feat-cons}
    \showconsensus{bottom}
\end{texshade}
```

1.5.4 Diversity mode

Contrary to the above described modes this shading style displays sequence differences. Thus, it is most suitable for comparing very similar sequences, e.g. species variants of a protein.

One sequence is used as consensus. Matching residues in other sequences are blanked out, mismatches are shown in lowercase. AQP2 species variants

	80	90	100	
Bos taurus	SFLRAVFYVA	AAQLİGAVAC	GAALLHEITPPA	ERG
Canis familiaris			h	
Dugong dugong	<mark>t</mark> l.		id	
Equus caballus	<mark>a</mark>		<mark>d</mark> .	r
Elephas maximus	<mark>t</mark> l.		d.	

Code:

```
\begin{texshade}{AQP2spec.ALN}
   \shadingmode{diverse}
   \setends{1}{77..109}
   \featureslarge
   \feature{top}{1}{77..109}{}AQP2 species variants}
   \namesrm\namessl
   \hidenumbering\showruler{top}{1}
   \shownames{left}
   \nameseq{1}{Bos taurus}
   \nameseq{2}{Canis familiaris}
   \nameseq{3}{Dugong dugong}
   \nameseq{4}{Equus caballus}
   \nameseq{5}{Elephas maximus}
   \frameblock{1}{82..82,106..106}{Red[1pt]}
```

1.5.5 Functionality modes

Displaying functional peptide similarities is one of T_EX shade's strong capabilities. Six functional shading modes are predefined; further user specific modes can easily be created. The examples may not look very impressive when printed in grayscale. Enjoy them on your screen or use color printouts. As mentioned before, all colors can be changed to others or to grays without restrictions (see chapter 5).

charge: residues which are charged at physiological pH (7.4) are shaded if their number at a position is higher than the threshold

AQP1.PRO	GLGI <mark>E</mark> IIGTLQLVLCVLATT <mark>D</mark> R. <mark>RR</mark> DLGGSAPL	170
AQP2.PRO	AVTV <mark>E</mark> LFLTMQLVLCIFAST <mark>D</mark> E. <mark>RR</mark> G <mark>D</mark> NLGSPAL	162
AQP3.PRO	GFFDQFIGTAALIVCVLAIV <mark>D</mark> PYNNPVPRGLEAF	186
AQP4.PRO	GLLV <mark>E</mark> LIITFQLVFTIFASC <mark>D</mark> S. <mark>KR</mark> T <mark>D</mark> VTGSVAL	191
AQP5.PRO	AMVV <mark>E</mark> LILTFQLALCIFSST <mark>D</mark> S. <mark>RR</mark> TSPVGSPAL	163

```
X acidic (-)
X basic (+)
```

```
\begin{texshade}{AQPpro.MSF}
    \shadingmode[charge]{functional}
    \setends{1}{138..170}
    \feature{top}{3}{153..165}{bar[-50,50]:-50,-45,%
        -40,-30,-20,-10,0,10,20,30,40,45,50}{}
    \feature{top}{3}{167..186}{color:5,10,15,20,25,30,35,%
        40,45,50,55,60,65,70,75,80,85,90,95,100[ColdHot]}{}
    \showlegend
\end{texshade}
```

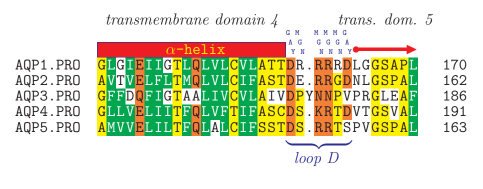
hydropathy: discrimination between acidic and basic, polar uncharged and hydrophobic nonpolar residues

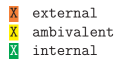
	lowercased	tinted	
	\sim		
AQP1.PRO	GLGI <mark>E</mark> IIG <mark>T</mark> LqlvlcVI	LA <mark>TT</mark> DR . RRRDL <mark>GGS</mark> APL	170
AQP2.PRO	AVTVELFLTMqlvlcIF	FA <mark>ST</mark> DE.RRGD <mark>NLGS</mark> PAL	162
AQP3.PRO	<mark>G</mark> FFDQFIG <mark>T</mark> AalivcVI	_AIVDPYNNPVPR <mark>G</mark> LEAF	186
AQP4.PRO	GLLVELII <mark>T</mark> FqlvftIF		191
AQP5.PRO	AMVVELILTFqlalcIF	F <mark>SST</mark> DS.RRTSP <mark>VGS</mark> PAL	163
	V pridic ()		
	X acidic (-)		
	X basic (+)		
	X polar uncharged		
	X hydrophobic nonpo	olar	
Cod	0.		
COC	IC.		

21

```
\begin{texshade}{AQPpro.MSF}
   \shadingmode[hydropathy]{functional}
   \feature{top}{1}{158..163}{brace}{tinted}
   \tintblock{1}{158..163}
   \feature{top}{1}{QLVLC}{brace}{lowercased}
   \lowerblock{1}{QLVLC}
   \setends{1}{138..170}
   \showlegend
\end{texshade}
```

structure: displays the potential localization within the tertiary structure of the protein





Code:

chemical: residues are shaded due to chemical properties of their functional groups

AQP1.PRO	GLGIEIIGTLQLVLCVLATTDR.RRRDLGGSAPL	170
AQP2.PRO	AVTVELFLTMQLVLCIFASTDE.RRGDNLGSPAL	162
AQP3.PRO	GFFDQFIGTAALIVCVLAIVDPYNNPVPRGLEAF	186
AQP4.PRO	GLLVELIITFQLVFTIFASCDS.KRTDVTGSVAL	191
AQP5.PRO	AMVVELILTFQLALCIFSSTDS.RRTSPVGSPAL	163
	<pre>X acidic (-) X aliphatic x aliphatic (small) X amide X aromatic X basic (+) X hydroxyl X imino X sulfur</pre>	

Code:

```
\begin{texshade}{AQPpro.MSF}
    \shadingmode[chemical]{functional}
    \setends{1}{138..170}
    \showlegend
\end{texshade}
```

With $\$ and all residues the threshold is ignored and all residues are shaded due to their group assignment. This is *not* identical to a threshold of 0% where only the majority group would be shaded. See the difference:

AQP1.PRO	GLGIEIIGTLQLVLCVLATTDR.RRRDLGG <mark>S</mark> APL	170
AQP2.PRO	AVTVELFLTMQLVLCIFASTDE.RRGDNLGSPAL	162
AQP3.PRO	GFFDQFIGTAALIVCVLAIVDPYNNPVPRGLEAF	186
AQP4.PRO	GLLVELII <mark>TFQ</mark> LVF <mark>T</mark> IFA <mark>SCDS</mark> .KRTDVTGSVAL	191
AQP5.PRO	A <mark>M</mark> VVELILTFQLALCIFSSTDS.RRTSPVG <mark>SP</mark> AL	163

Code:

```
\begin{texshade}{AQPpro.MSF}
   \shadingmode[chemical]{functional}
   \setends{1}{138..170}
   \shadeallresidues
\end{texshade}
```

rasmol: similar to [chemical] but with shading following the rasmol color scheme

	∠ 🗅	
	1000 1000	
AQP1.PRO	GLGIEIIGTLQLVLCVLATTDR.RRRDLGG <mark>S</mark> APL	170
AQP2.PRO	AVTVELFLTMQLVLCIFASTDE.RRGDNLGSPAL	162
AQP3.PRO	GFFDQFIGTAALIVCVLAIVDPYNNPVPRGLEAF	186
AQP4.PRO	GLLVELIITFQLVFTIFASCDS.KRTDVTGSVAL	191
AQP5.PRO	AMVVELILTFQLALCIFSSTDS.RRTSPVGSPAL	163

X	Asp,	Glu	
Х	Arg,	Lys,	His
Х	Phe,	Tyr,	Trp
Х	Ala,	Gly	
X	Cys,	Met	
X	Ser,	Thr	
X	Asn,	Gln	
Χ	Leu,	Val,	Ile
Х	Pro		

```
\begin{texshade}{AQPpro.MSF}
   \shadingmode[rasmol]{functional}
   \setends{1}{138..170}
   \showruler{bottom}{1}
   \rulersteps{1}
   \namerulerpos{150}{site A[Red]}
   \namerulerpos{155}{site B[Green]}
   \shadeallresidues
   \showlegend
\end{texshade}
```

standard area: this shading displays the differences in the surface area of the different amino acid's sidechains

AQP1.PRO	GLGIEIIGTLQLVLCVLATTDR.RRRDLGGSAPL	170
AQP2.PRO	AVTVELFLTMQLVLCIFASTDE.RRGDNLGSPAL	162
AQP3.PRO	GFFDQFIGTAALIVCVLAIVDPYNNPVPRGLEAF	186
AQP4.PRO	GLLVELIITFQLVFTIFASCDS.KRTDVTGSVAL	191
AQP5.PRO	AMVVELILTFQLALCIFSSTDS.RRTSPVGSPAL	163

```
Х
   88.1 (G); Standard sidechain area (Å^2)
  118.2 (A); 129.8 (S)
Х
X
  146.1 (C); 146.8 (P)
Х
  152.5 (T); 158.7 (D); 164.5 (V); 165.5 (N)
Х
  181.0 (I); 186.2 (E)
Х
  193.1 (L); 193.2 (Q); 202.5 (H); 203.3 (M)
Х
  222.8 (F); 225.8 (K)
Х
  238.8 (Y)
  256.0 (R); 266.2 (W)
Х
```

```
\begin{texshade}{AQPpro.MSF}
   \shadingmode[standard area]{functional}
   \setends{1}{138..170}
   \showlegend
   \shadeallresidues
\end{texshade}
```

accessible area: here, the surface area which can be accessed by solvent molecules is used as a basis for shading; low accessibility means hydrophobic (i. e. strongly buried residues), whereas highly accessible sidechains are hydrophilic (compare to hydropathy and structure)

	membr.	loop	membr.	
	_0000000000000000000			
AQP1.PRO	GLGIEIIGTLQLVLCVLATTD	R.RRI	R <mark>DLGGSA</mark> PL	170
AQP2.PRO	AVTVELFLTMQLVLCIFASTD	E.RR	GDNLGSPAL	162
AQP3.PRO	GFFDQFIGTAALIVCVLAIVD			186
AQP4.PRO	GLLVELIITFQLVFTIFASC D	S.KR	F<mark>DVT</mark>GSVAL	191
AQP5.PRO	AMVVELILTFQLALCIFSSTD	S.RR	FS<mark>PVG</mark>S<mark>PAL</mark>	163

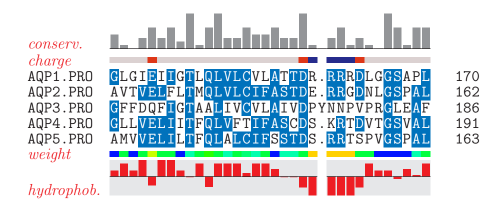
13.9 (C); Accessible sidechain area (Å²) 23.0 (I); 23.5 (V); 25.2 (G) Х Х 28.7 (F); 29.0 (L); 30.5 (M); 31.5 (A) Х 41.7 (W); 44.2 (S); 46.0 (T); 46.7 (H) Х 53.7 (P) Х 59.1 (Y); 60.9 (D); 62.2 (N) Х 72.3 (E); 74.0 (Q) Х 93.8 (R) Х 110.3 (K)

Code:

```
\begin{texshade}{AQPpro.MSF}
   \shadingmode[accessible area]{functional}
   \setends{1}{138..170}
   \showlegend
   \feature{top}{1}{138..157,164..170}{helix}{membr.}
   \feature{top}{1}{158..163}{---}{loop}
   \featurerule{1mm}
   \shadeallresidues
\end{texshade}
```

1.6 Bar graphs and color scales

Amino acid properties, such as hydrophobicity, molecular weight, or charge can be shown as bar graphs or color scales along the alignment. Further, the degree of protein sequence conservation can be indicated. As an example, in the following aquaporin alignment plots of residue conservation (bars, top), are shown as well as properties of the AQP1 sequence: charge (scale, top), molecular weight are shown (scale, bottom), and hydrophobicity (bars, bottom).



```
\begin{texshade}{AQPpro.MSF}
   \setends{1}{138..170}
   \feature{ttop}{1}{138..170}{bar:conservation}{}
   \showfeaturestylename{ttop}{conserv.}
   \ttopspace{-\baselineskip}
   feature{top}{1}{138..170}{color:charge}{}
   \showfeaturestylename{top}{charge}
   \feature{bottom}{1}{138..170}{color:molweight[ColdHot]}{}
   \showfeaturestylename{bottom}{weight}
   \bbottomspace{-\baselineskip}
   \feature{bbottom}{1}{138..170}{bar:hydrophobicity[Red,Gray10]}{}
   \showfeaturestylename{bbottom}{hydrophob.}
   \bargraphstretch{3}
   \featurestylenamescolor{Red}
   \featurestylenamesrm \featurestylenamesit
   \hideconsensus
\end{texshade}
```

The degree of similarity and identity between all sequences in the alignment can be shown in a table using \similaritytable (4.10.2) outside the texshade environment:

	AQP1.PRO	AQP2.PRO	AQP3.PRO	AQP4.PRO	AQP5.PRO	
AQP1.PRO		69.6	42.4	63.6	63.6	
AQP2.PRO	48.4		36.3	66.6	75.7	similarity
AQP3.PRO	33.3	21.2		33.3	39.3	lila
AQP4.PRO	45.4	54.5	24.2		75.7	sin
AQP5.PRO	42.4	66.6	24.2	57.5		8
	% ide	entity				

1.7 Secondary structures

Predicted protein secondary structures in the DSSP, STRIDE PHD or HMMTOP file format can be included and displayed in the alignment. As an example, the following few commands show an aquaporin alignment with the PHD topology data for aquaporin type 1 (top sequence).

```
\begin{texshade}{AQPpro.MSF}
    \shadingmode[allmatchspecial]{similar}
    \includePHDtopo{1}{AQP1.phd}
\end{texshade}
```

Abbr.: int.	– internal; $ext.$ – external; TM – transmembrane de	omain
AQP1.PRO AQP2.PRO AQP3.PRO AQP4.PRO AQP5.PRO consensus	MASEIKKKLFW MWELRSIAFS MNRCGEMLHIRYRLL MSDGAAARRWGKCGPPCSRESIMVAFKGVWTQAFW MKKEVCSLAFF ! *** * ***	11 10 15 35 11

int. A

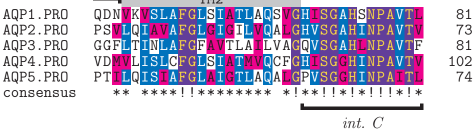
ext. B TM1 RAVVAEFLAMTLFVFISIGSALGFNYPLERNQTLV RAVLAEFLATLLFVFFGLGSALQWA...SS...P RQALAECLGTLILVMFGCGSVAQVVLSRGTH....

AQP3.PRO AQP4.PRO AQP5.PRO consensus

AQP1.PRO

AQP2.PRO

		-
.PRO	KAVTAEFLAMLIFVLLSVGSTINWGGSENPLP	67
.PRO	<mark>KAVFAEFLATLIFVFFGLGSAL</mark> KWPSAL	39
ensus	**** * ***** **** !**** *	
	int. A	
	ext. B	
	TM2	



ext. D

TM/

46

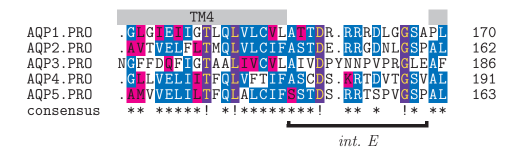
38

46

	TM3	
AQP1.PRO	GLLLSCQISILRAVMYIIAQCVGAIVASAILSGI.	115
AQP2.PRO	ACLVGCHVSFLRAAFYVAAQLLGAVAGAAILHEI.	107
AQP3.PRO	AMCFLAREPW <mark>IK</mark> LPIYTLAQTLGA <mark>FLGAGIVF</mark> GLY	116
AQP4.PRO	AMVCTRKISIAKSVFYITAQCLGAIIGAGILYLV.	136
AQP5.PRO	ALLIGNQISLIRAVFYVAAQLVGAIAGAGILYWL.	108
consensus	<u>****</u> <u>********* ii *ii*</u> *********	

ext. D

AQP1.PRO		137
AQP2.PRO		129
AQP3.PRO	YDAIWAFAGNE <mark>L</mark> V <mark>V</mark> S <mark>G</mark> PN <mark>G</mark> TA <mark>GI</mark> FATYPS <mark>G</mark> HLDMV	151
AQP4.PRO	TPPS <mark>V</mark> VGGLGVTTVHGNLTAGH	158
AQP5.PRO		130
consensus	*** * * ***** * * !	

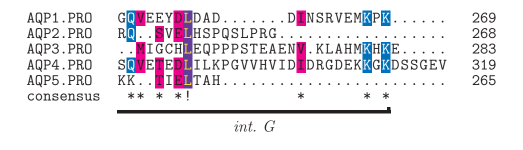


ext.	F
------	---

	1M5	
AQP1.PRO	AIGLSVALGHLLAIDYTGCGINPARSFGSAVLTR.	204
AQP2.PRO	SIGFSVTLGHLLGIYFTGCSMNPARSLAPAVVTG.	196
AQP3.PRO	TVGLVVLVIGTSMGFNSGYAVNPARDFGPRLFTAL	221
AQP4.PRO	AIGFSVA <mark>IGHLFAINY</mark> TGASMNPARSFGPAVIMG.	225
AQP5.PRO	SIGLSVTLGHLVGIYFTGCSMNPARSFGPAVVMN.	197
consensus	<u>**i**i</u> **** <u>**</u> ***********************	

	ext. F	
AQP1.PRO	TM6 .N <mark>F</mark> S.NHWIFWVGPFIGSALAVLIYDF	229
AQP2.PRO AQP3.PRO	.KFD.DHWVFWIGPLVGATIGSLLYNY AGWGSEVFTTGQNWWWVPIVSPLLGSIGGVFVY	221 254
AQP4.PRO AQP5.PRO	.NWE.NHWIYWVGPIIGAVLAGALYEY .RFSPSHWVFWVGPIVGAMLAAILYFY	250 223
consensus	* * * * * * * * * * * * * * * * * *	
AQP1.PRO AQP2.PRO	ILAPRSSDFTDRMKVWTS LLFPSAKSLQERLAVLKG.LEPDTDWEEREVRR	247 253
AQP3.PRO AQP4.PRO	QL	256 284
AQP5.PRO consensus	LLFPSSLSLHDRVAVVKGTYEPEEDWEDHREER **** * *** * ** * * *	256

int. G



AQP1.PRO		269
AQP2.PRO	.SKA	271
AQP3.PRO	QI	285
AQP4.PRO	LSSV	323
AQP5.PRO		265
consensus		

1.8 Sequence fingerprints

To gain a quick overview of sequence similarities or properties the **\fingerprint** command has been implemented. It can depict the complete sequence in one single line. The residues are presented as colored vertical lines. The implementation of this kind of output was inspired by a publication by KAI-UWE FRÖHLICH [6].

	TM	TM TI	$M \qquad TM \ TM$	TM
AQP1.PRO				
AQP2.PRO				
AQP3.PRO				
AQP4.PRO				
AQP5.PRO				

non	conserved
sim	ilar
\geq 50)% conserved
all	match

Code:

The higher the similarity the darker the vertical lines. In this overview it becomes obvious that the transmembrane regions of the aquaporin isoforms are most conserved.

A fingerprint of charge distribution on different aquaporins is shown. below. Sequence gaps can be left blank (example above) or drawn as lines between the sequence blocks.

AQP1.PRO				
AQP2.PRO				- - -
AQP3.PRO			-	
AQP4.PRO				
AQP5.PRO	├ 			

Code:

```
\begin{texshade}{AQPpro.MSF}
    \shadingmode[charge]{functional}
    \shadeallresidues
    \fingerprint{360}
    \gapchar{rule}
    \showlegend
\end{texshade}
```

1.9 Sequence logos

Sequence logos represent the information content of the aligned sequences at a position in bit (max. 2 bit for DNA, i. e. $\log_2 4$, and 4.322 bit for proteins, i. e. $\log_2 20$) and the relative frequency of a base or amino acid at this position [7]. Thus, more information is contained in logos than in a standard consensus sequence. The example below shows a DNA sequence alignment with the logo on the top.

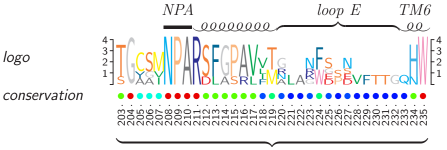
It must be remarked that a logo from only five sequences does not produce meaningful results - it rather illustrates the technique.



Code:

```
\begin{texshade}{AQPDNA.MSF}
   \setends{1}{414..443}
   \showsequencelogo{top}
\end{texshade}
```

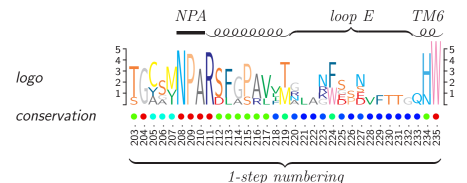
Next, only the logo of a protein alignment is displayed plus the degree of sequence conservation as a color scale in the consensus line. Note, that the full functionality of the feature lines remains.



1-step numbering

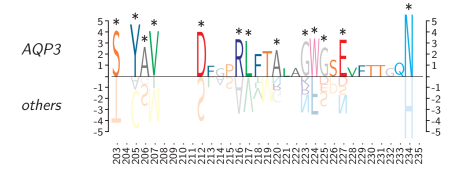
```
\begin{texshade}{AQPpro.MSF}
   \setends{AQP3.PR0}{203..235}
   \showsequencelogo{top} \showlogoscale{leftright}
   \hideseqs
   \residuesperline*{33}
   \defconsensus{{$\bullet$}}{{$\bullet$}}{{$\bullet$}}
   \showconsensus{ColdHot]{bottom}
   \nameconsensus{conservation} \namessf\namessl
    \showruler{bottom}{AQP3.PR0} \rulersteps{1}
   \feature{top}{AQP3.PR0}{208..210}{---}{NPA}
   \feature{top}{AQP3.PR0}{211..219}{helix}{}
   \feature{top}{AQP3.PR0}{220..232}{brace}{loop E}
   \feature{top}{AQP3.PR0}{203..235}{brace}{1-step numbering}
   \end{texshade}
```

The same logo is shown below but with frequency correction turned on (\dofrequencycorrection), see p.54. This takes into account the difference between the amino acid distribution in the alignment and the equal distribution of 5% for each residue.



1.10 Subfamily logos

The following output is derived from the calculation of a subfamily logo [14]. Such logos display relevant deviations of a subfamily compared to the remaining set of sequences. Here, typical residues of AQP3 are shown (upright) which deviate from the remaining four aquaporins of this alignment (upside-down). The output can be directly compared to the sequence logo above, which displays the same section of the alignment. Note, that five sequences are far too few to obtain meaningful results with this method. This is just to illustrate the approach.



Code:

```
\begin{texshade}{AQPpro.MSF}
   \setends{AQP3.PR0}{203..235}
   \residuesperline*{33}
   \setsubfamily{3}
   \showsubfamilylogo{top} \showlogoscale{leftright}
   \namesubfamilylogo[others]{AQP3}
   \namessf \namessl
   \showruler{bottom}{AQP3.PR0} \rulersteps{1}
   \hideseqs
   \hideconsensus
   \dofrequencycorrection
\end{texshade}
```

1.11 Customization of the alignment output

Extensive possibilities are given to the user to customize the final output of an alignment. Thus, all parameters defining the appearance of letters can be changed individually for sequence residues, names and numbering or the describing feature texts. Additional manual shading can be applied to any region or block of residues. Sequences are easily re-ordered, separated, hidden or blanked out without recalculation of the entire alignment; sections of the alignment can also be shown. Numbering and rulers can be displayed and set to any value. A powerful tool is the **\feature** command which allows one to label stretches of residues with bars, arrows, braces or any fill character and describing text. Legends are set automatically if desired, but user commands are also provided to build individual legends.

2 Format of alignment input files

T_EX<mark>shade</mark> can handle two common alignment input formats, i.e. the MSF format (multiple sequence format) and the ALN format (alignment format). The MSF format is used by PILEUP of the Unix GCG sequence analysis package²¹. Files in the ALN format are produced by CLUSTAL which is available for free for Unix, DOS and Macintosh. Further, upon request, the FASTA format is supported since version 1.6. In addition to the mentioned software many alignment programs have export filters for the MSF, ALN or FASTA format, e.g. MACAW produces ALN files. If you are not sure whether your favorite sequence aligner produces one of the required formats compare its output to the following examples. T_FXshade determines the format from the internal file structure, thus extensions like MSF, ALN or FASTA are not required. If you can choose the alignment format MSF is recommended, because this format gives information about the sequence type, i.e. peptide or nucleotide sequences, and length (for the correct setting of gaps at the sequence end).

2.1 The MSF file format

Files of this type are divided into a header section and the multiple sequence alignment. The header may contain the following components:

File Type: (optional) The first header line reads for nucleic acids alignments !!NA_MULTIPLE_ALIGNMENT 1.0 and for amino acid sequences !!AA_MULTIPLE_ALIGNMENT 1.0 (all uppercase).

Description: (optional) Informative text describing what is in the file.

Dividing line: (required!) Must include the following attributes:

- MSF: Displays the number of bases or residues in the multiple sequence alignment.
- Type: Displays the sequence type, 'P' for a peptide and 'N' for a nucleotide alignment.
- Checksum: Displays an integer value that characterizes the contents of the file.

²¹For a description see http://gene.md.huji.ac.il/Computer/GCG9doc

- ... The two periods act as a divider between the descriptive information and the following sequence information.
- Name/Weight: (required!) Must include the name of each sequence included in the alignment, as well as its length, checksum and weight.
- **Two slashes** (//): (required!) This separating line divides the name/weight information from the sequence alignment

The alignment section consists of sequence blocks divided by an empty line. Each sequence line starts out with the sequence name. An example file is shown here:

AQP.MSF	MSF: 87 Type:	P May 1st,	1998	Check: 2	2586
Name: AQP	1.PRO Len: 66	Check: 13	67	Weight:	1.00
Name: AQP	2.PRO Len: 58	Check: 21	76	Weight:	1.00
Name: AQP	3.PRO Len: 83	Check: 18	93	Weight:	1.00
Name: AQP	4.PRO Len: 63	Check: 37	37	Weight:	1.00
Name: AQP	5.PRO Len: 59	Check: 34	13	Weight:	1.00
11				-	
	1				45
AQP1.PRO	MAS		EIKK	KLFWRAV	/AEFLAM
AQP2.PRO	MW		ELRS	IAFSRAVI	LAEFLAT
AQP3.PRO	MNRCG	EMLHIR	YR	LLRQAI	LAECLGT
AQP4.PRO	MSDGAAARRWGKCG	PPCSRESIMVA	FKGVWT	QAFWKAVI	TAEFLAM
AQP5.PRO	MK		.KEVCS	LAFFKAVH	FAEFLAT
	45				87
AQP1.PRO	TLFVFISIGSALGF	NYPLERNQTLV	QDNVKV	SLAFGLSI	LATL
AQP2.PRO	LLFVFFGLGSALQW	AP	PSVLQI	AVAFGLG	IGIL
AQP3.PRO	LILVMFGCGSVAQV	VLSRGTHGGF.	LTI	NLAFGFA	/TLA
AQP4.PRO	LIFVLLSVGSTINW	GGSENPLP	VDMVLI	SLCFGLSI	LATM
AQP5.PRO	LIFVFFGLGSALKW	PSAL	PTILQI	SIAFGLAI	IGTL

TEXSTAGE extracts only the information from the file it really needs. So, do not mind all the checksums listed in the file—TEXSTAGE does not either. The same is true for Weight. Required are the string MSF: for the identification of the file format and Type: for the determination of the sequence type (both in the dividing line), further all Name: definitions and finally //. The MSF format allows one to comment out sequences. This is done by putting an exclamation point directly infront of the respective Name. These sequences are neither displayed nor used for the calculation of the consensus. This works for T_EX shade, too. To comment out sequences without changing the input file use the T_EX shade command \sillseq{seqref} (4.5.3).

Name: AQP !Name: AQ !Name: AQP Name: AQP	MSF: 87 Type: P May 1st, 1998 Check: 2586 1.PRO Len: 66 Check: 1367 Weight: 1.00 P2.PRO Len: 58 Check: 2176 Weight: 1.00 P3.PRO Len: 83 Check: 1893 Weight: 1.00 4.PRO Len: 63 Check: 3737 Weight: 1.00 5.PRO Len: 59 Check: 3413 Weight: 1.00
//	
	1 45
AQP1.PRO	MASEIKKKLFWRAVVAEFLAM
AQP2.PRO	MWELRSIAFSRAVLAEFLAT
AQP3.PRO	MLLRQALAECLGT
AQP4.PRO	MSDGAAARRWGKCGPPCSRESIMVAFKGVWTQAFWKAVTAEFLAM
AQP5.PRO	MKKEVCSLAFFKAVFAEFLAT
	45 87
AQP1.PRO	TLFVFISIGSALGFNYPLERNQTLVQDNVKVSLAFGLSIATL
AQP2.PRO	LLFVFFGLGSALQWASSPPSVLQIAVAFGLGIGIL
AQP3.PRO	LILVMFGCGSVAQVVLSRGTHGGFLTINLAFGFAVTLA
AQP4.PRO	LIFVLLSVGSTINWGGSENPLPVDMVLISLCFGLSIATM
AQP5.PRO	LIFVFFGLGSALKWPSALPTILQISIAFGLAIGTL
ΑΨΓΟΤΓΙΟ	TTLALAFOOVPUMLOVPLITFATOTM, AFVIALF

The sequence lengths given after Len: are not used by T_EX shade. Due to the fact that most alignment programms calculate the sequence length by summing up residues and additionally gaps which is not really correct. In order to have the sequence break right after the last residue without printing further gap symbols T_EX shade counts the number of residues by itself. You can also use the command seqlength in the T_EX shade environment to set the values manually if you do not trust a machine.

2.2 The ALN file format

ALN files are quite similar to the above described MSF files. They simply lack a defined header section. Nevertheless, describing text is allowed before the alignment part. T_EX shade determines the number of sequences and their names from the last sequence block—so, no further text lines are allowed after this block! Due to a lacking

declaration in the file the sequence type has to be set in the texshade environment by $seqtype{\langle type \rangle}$ with 'P' for peptide and 'N' for nucleotide sequences; for the example below: $seqtype{P}$. If no seqtype command is used T_FXshade assumes a nucleotide sequence.

profalign May 1st, 1998, 16:58 of AQPpro.MSF{} Muliple alignment parameter: Gap Penalty (fixed): 10.00 Gap Penalty (varying): .05 Gap separation penalty range: 8 Percent. identity for delay: 0% List of hydrophilic residue: GPSNDQEKRH Protein Weight Matrix: blosom 10 20 30 40 AQP1.PRO MAS.....EIKKKLFWRAVVAEFLAM AQP2.PRO MW.....ELRSIAFSRAVLAEFLAT AQP3.PRO M.....NRCG....EMLHIRYR....LLRQALAECLGT AQP4.PRO MSDGAAARRWGKCGPPCSRESIMVAFKGVWTQAFWKAVTAEFLAM AQP5.PRO MK.....KEVCSLAFFKAVFAEFLAT * ** * AQP1.PRO TLFVFISIGSALGFNYPLERNQTLVQDNVKVSLAFGLSIATL AQP2.PRO LLFVFFGLGSALQWA...SS....PPSVLQIAVAFGLGIGIL AQP3.PRO LILVMFGCGSVAQVVLSRGTHGGF....LTINLAFGFAVTLA AQP4.PRO LIFVLLSVGSTINWG...GSENPLPVDMVLISLCFGLSIATM AQP5.PRO LIFVFFGLGSALKWP....SA....LPTILQISIAFGLAIGTL .. * .** **

The minimal contents of an ALN file are shown below; this is fully sufficient. Many sequence alignment programs can produce such an output. Have a look at **seqpup** by DON GILBERT if you need a comprehensive conversion program²².

²²Sorry, seqpup is much more!

```
AQP1.PR0MAS.EIKKKLFWRAVVAEFLAMAQP2.PR0MW.ELRSIAFSRAVLAEFLATAQP3.PR0M..NRCG.EMLHIRYR.AQP4.PR0MSDGAAARRWGKCGPPCSRESIMVAFKGVWTQAFWKAVTAEFLAMAQP5.PR0MK..KEVCSLAFFKAVFAEFLATAQP1.PR0TLFVFISIGSALGFNYPLERNQTLVQDNVKVSLAFGLSIATLAQP2.PR0LLFVFFGLGSALQWA.AQP3.PR0LILVMFGCGSVAQVVLSRGTHGGF.AQP4.PR0LIFVFFGLGSALKWP.AQP5.PR0LIFVFFGLGSALKWP.
```

2.3 The FASTA file format

In FASTA files each sequence is led by a single description line starting with a '>'. T_EX **shade** uses the first word delimited by the leading '>' and a space as the sequence name. If no descriptive text is present T_EX **shade** generates a sequence name consisting of 'seq' plus a consecutive number. The lines following the description line contain the sequence.

```
>AQP1.PR0
MAS.....EIKKKLFWRAVVAEFLAM
TLFVFISIGSALGFNYPLERNQTLVQDNVKVSLAFGLSIATL
>AQP2.PR0
MW.....ELRSIAFSRAVLAEFLAT
LLFVFFGLGSALQWA...SS...PPSVLQIAVAFGLGIGIL
>AQP3.PR0
M....NRCG...EMLHIRYR...LLRQALAECLGT
LILVMFGCGSVAQVVLSRGTHGGF...LTINLAFGFAVTLA
>AQP4.PR0
MSDGAAARRWGKCGPPCSRESIMVAFKGVWTQAFWKAVTAEFLAM
LIFVLLSVGSTINWG...GSENPLPVDMVLISLCFGLSIATM
>AQP5.PR0
MK....KEVCSLAFFKAVFAEFLAT
LIFVFFGLGSALKWP...SA...LPTILQISIAFGLAIGTL
```

3 Use of a T_EXshade parameter file

Using predefined parameter files for repeatedly occuring situations can save a lot of typing and makes the output throughout the publication or presentation more consistent. Further, such files are an easy way to exchange self-defined shading modes or new color schemes (i. e. for a satisfying grayscale output) with other users. If you have created a parameter file, which you think is of interest for others, please submit it to me²³ as an e-mail attachment together with a short description. I will take care of those files and post them—with a reference to the author—together with the next T_EX**shade** distribution to make them available for all interested users.

No special file format is required for parameter files. T_EX_{shade} simply calls the file using the \input command right after resetting all parameters to default. An example parameter file is present containing the standard parameters of T_EX_{shade} called texshade.def. This file can be changed freely and can be used as a template for the creation of personal parameter files.

Five steps are executed by $T_E X shade$ when processing the texshade environment:

$\begin{texshade}[\langle parameterfile \rangle]{\langle alignmentfile \rangle}$

- 1. Analysis of the *(alignmentfile)*; determination of the number of sequences and sequence names
- 2. Setting parameters to default
- 3. Setting parameters to the definitions of the $\langle parameterfile \rangle$, if existent
- 4. Execution of further $T_E X$ **shade** commands within the evironment, if existent

\end{texshade}

5. Loading and setting the alignment on a line by line basis

 $^{^{23} \}tt ebeitz@pharmazie.uni-kiel.de$

4 texshade user commands

The T_EX shade package must be loaded by the \usepackage command in the document header section.

 $\ [\langle option \rangle] \{ texshade \}$

Then, the texshade environment is ready to use as described in 1.4. See also section 3 for a description of the optional parameter file. All other commands provided by T_EX shade (except \shadebox [4.7], \molweight and \charge [4.10.1], and \percentsimilarity, \percentidentity and \similaritytable [4.10.2]) must be used within the texshade environment.

4.1 Using predefined shading modes

If no \shadingmode command is given in the texshade environment the default shading mode (*identical*, see 1.5.1) is active. For the selection of one of the other predefined shading modes the following command is provided.

You can choose from four shading modes and declare one option which depends on the selected mode.

See 1.5.1 for examples. Use the option allmatchspecial to shade positions with a special color where all residues are identical. Or use a percentage number (0-100) as an option to set an additional threshold for highly conserved residues, e.g. \shadingmode[90]{identical}. \allmatchspecial can also be used as a command with or without an optional parameter for setting the high conservation threshold. As both, option or command, allmatchspecial is only active in the *identical* and *similar* shading modes.

 conserved) residues can be set individually plus the letter case (lower or upper) or any character can be chosen:

```
\label{eq:conserved} $$ \conservedresidues{$$ (res.col.)}{$$ (shad.col.)}{$$ (case)}{$$ (style)} $$ allmatchresidues{$$ (res.col.)}{$$ (shad.col.)}{$$ (case)}{$$ (style)} $$
```

For how to handle colors for the foreground $\langle res.col. \rangle$ and the background $\langle shad.col. \rangle$ see section 5. The third parameter $\langle case \rangle$ tells T_EX**shade** to print the corresponding residue as a lowercase or an uppercase letter or even to print any other character. Finally, the $\langle style \rangle$ parameter tells T_EX**shade** which shape to use for the letters. Use one of the following styles for $\langle style \rangle$.

$\langle style \rangle$	effect
bf	bold face series
md	normal series
up	upright shape (normal shape)
it	italics shape
sl	slanted shape
rm	modern roman family
sf	sans serif family
tt	typewriter family

In order to change only some of the parameters it is sufficient to declare these and use empty braces for the others. Examples:

\conservedresidues{White}{Blue}{upper}{bf}: the conserved residues are printed as bold face white uppercase letters on blue.

 $\mbox{nomatchresidues}}{}:$ instead of the non matching residues a '•' is printed. The colors and style are not changed. Note the double curly braces which make T_EX shade interpret this complex symbol description as one single character.

Once a set of shading colors is redefined, it can be saved using $defshadingcolors{\langle name \rangle}$ for later use in the document (see 4.6.1).

2. $\ \ [\langle all match special / number \rangle] \{ similar \}$

See 1.5.2 for an example output and an explanation of the shading. In addition to the described commands for changing shading colors this shading mode provides the command \similarresidues. Use it in analogy to the commands above.

How does T_EX **shade** know which residues are considered to be similar? These definitions are set by two command couples, i.e. \pepsims,\pepgroups for peptides and \DNAsims,\DNAgroups for nucleotides. With \pepsims and \DNAsims residues are defined which are similar to the consensus residue. Examples:

\pepsims{S}{TA} If a serine is the consensus residue then all threenins and alanines at this position are shaded in the color for similars. This definition does *not* imply that threenine and alanine are similar to each other! This becomes obvious when you inspect the next definition:

 $pepsims{T}{S}$ Serine but not alanine is declared to be similar to threenine.

What happens if there is no consensus residue? How does T_EX shade decide if a group of similars is greater than the threshold? For this groups are pre-defined:

\pepgroups{FYW,ILVM,RK,DE,GA,ST,NQ} This command allows one to set up to nine groups of similars, separated by commas. Each residue can belong to only one group. If one residue is assigned to several groups only the last assignment is carried out.

\DNAgroups{GAR,CTY} This command is used in analogy to the amino acid groups. Here, two ambiguity codes ('R' for purine base and 'Y' for pyrimidine base) are assigned in addition.

Residues which do not appear in any of the four commands are considered not to belong to a group. The default settings for similars are listed below:

\pepgroups{FYW,ILVM,RK,DE,GA,ST,NQ}

\pepsims{F}{YW} % Y and W are similar to F \pepsims{Y}{WF} % W and F are similar to Y \pepsims{W}{YF} % Y and F are similar to W

```
\pepsims{I}{LVM} % L, V and M are similar to I
\pepsims{L}{VMI} % V, M and I are similar to L
\pepsims{V}{MIL} % M, I and L are similar to V
pepsims{R}{KH}
                % K and H are similar to R
                % H and R are similar to K
pepsims{K}{HR}
pepsims{H}{RK}
                % R and K are similar to H
                % G and S are similar to A
pepsims{A}{GS}
pepsims{G}{A}
                % A (but not S) is similar to G
pepsims{S}{TA}
                \% T and A are similar to S
                % S (but not A) is similar to T
pepsims{T}{S}
pepsims{D}{EN}
                \% E and N (but not Q) are similar to D
\pepsims{E}{DQ}
                % D and Q (but not N) are similar to E
pepsims{N}{QD}
                % Q and D (but not E) are similar to N
                % N and E (but not D) are similar to Q
pepsims{Q}{NE}
\DNAgroups{GAR,CTY}
DNAsims{A}{GR}
                % G and R are similar to A
DNAsims{G}{AR}
                % A and R are similar to G
DNAsims{R}{AG}
                % A and G are similar to R
                % T and Y are similar to C
DNAsims{C}{TY}
                % C and Y are similar to T
DNAsims{T}{CY}
```


 $DNAsims{Y}{CT} % C and T are similar to Y$

Enter a $\langle filename \rangle$ to load the shading information from a T-Coffee score_ascii file (www.tcoffee.org); see example in 1.5.3. Make sure that the alignment file specified in the \texshade command and this shading file correspond to each other.

If you do not enter a $\langle filename \rangle$ here, a separate command $\includeTCoffee{\langle filename \rangle}$ must be used.

T-Coffee shading can also be used in the consensus p. 51 and in the feature lines, in particular color scales and bar plots p. 71, for the display of shading information.

1.5.4 depicts an example alignment. Choose the number or the name of the sequence $\langle seqref \rangle$ which will be treated as the consensus and to which the other sequences are compared. If no $\langle seqref \rangle$ is declared the first sequence is set as consensus ($\langle seqref \rangle = 1$).

Standard definitions for diverse mode are:

```
\nomatchresidues{Black}{White}{lower}{up}
\similarresidues{Black}{White}{lower}{up}
\conservedresidues{Black}{White}{{.}}{up}
\allmatchresidues{Black}{White}{{.}}{up}
\gapchar{-}
```

After calling \shadingmode{diverse} these commands can be used to redefine the diverse mode settings (mind the double curly braces around the dot-symbol!).

- 5. \shadingmode[$\langle type \rangle$]{functional} There are seven different functional shading modes available for peptide sequences; nucleotide sequences can not be shaded due to functional aspects. Five of TEXshade's functional modes correspond to the four 'alphabets' employed by KARLIN and GHANDOUR for peptide alignments [2] or by the rasmol software. Additional 'alphabets' to the standard 20-letter array of amino acids can highlight peptide similarities which were otherwise not visible. For the 'alphabet' definitions see below:
 - $\langle type \rangle$ = charge Acidic (D, E) and basic (H, K, R).
 - \$\langle type \rangle = hydropathy Acidic and basic (as above), polar uncharged (C, G, N, Q, S, T, Y) and hydrophobic nonpolar (A, F, I, L, M, P, V, W), see also KYTE and DOOLITTLE [3].
 - \$\langle type \rangle = structure External (D, E, H, K, N, Q, R), internal (F, I, L, M, V) and ambivalent (A, C, G, P, S, T, W, Y).
 - \$\langle type \rangle\$ = chemical Acidic (D, E), aliphatic (I, L, V), aliphatic (small) (A, G), amide (N, Q), aromatic (F, W, Y), basic (H, K, R), hydroxyl (S, T), imino (P) and sulfur (C, M).

⟨type⟩ = rasmol (D, E), (K, R, H), (F, Y, W), (A, G), (C, M), (S, T), (N, Q), (I, L, V), (P).

The two modes described below highlight sidechain sizes and hydrophobicity, respectively, according to ROSE *et al.* [4,5]. Standard area stands for the surface area of the residue in Å², i.e. it is a measure for the size of a residue's sidechain. The accessible area value (also in Å²) gives information about the size of the surface area which is accessible by solvent molecules within the folded protein. A very small area means that the residue is strongly buried and is thus very hydrophobic. Hydrophilic residues in turn possess large accessible areas due to their prefered location at the protein surface. Therefore, this kind of shading provides another method, in addition to hydropathy and structure, for the visualization of structural protein properties.

- $\langle type \rangle =$ standard area for the area values see legend of the alignment in 1.5.5
- $\langle type \rangle$ = accessible area for values see 1.5.5

If no $\langle type \rangle$ or an unknown $\langle type \rangle$ is designated as option all functional groups and shading colors are cleared. This is also achieved by the command \clearfuncgroups. With all groups cleared one can start to build new shading modes from scratch. How to do this is explained in the next section.

In order to exchange the colors but to keep the group definitions and descriptions the command \funcshadingstyle can be employed. Usage:

$$\label{eq:linearized_linearized$$

 $\langle residue \rangle$ is one representative of the whole amino acid group. The colors which are declared by the next four parameters are used for all residues in this group. $\langle case \rangle$ and $\langle style \rangle$ are as described for example in \nomatchresidues.

With \shadeallresidues the threshold is ignored and all residues are shaded due to their group assignment.

4.2 Creating new functional shading modes

The grouping of amino acids due to other properties can make sense as suggested by KARLIN and GHANDOUR [2], e.g. physical properties (molecular weight, shape), kinetic properties (reaction velocity, Michaelis-Menton constant), or structure (α -helices, β -sheets, turns). New amino acid groups are defined with the \funcgroup command. This command needs six parameters:

 $\label{eq:constraint} $$ \constraints \con$

 $\langle descr \rangle$ contains descriptive text which is displayed in the legend. The second parameter $\langle residues \rangle$ holds the amino acids to be grouped. The colors for the foreground and background are set with the following two parameters, the case and style is declared by the last parameters. The example below defines a functional group named 'acidic (-)' containing the amino acids aspartic and glutamic acid with white letters on a red background:

```
\funcgroup{acidic ($-$)}{DE}{White}{Red}{upper}{up}
```

For the usage of colors see section 5. Up to nine individual groups can be defined. New groups are simply added to the already existing groups, i.e. if an extension of the group definitions of an existing shading mode is desired there is no need to clear these groups und re-define them again. Just add the new groups with the \funcgroup command. To create completely new modes use the command \shadingmode{functional} without an option before setting the new groups. The new definitions are active only in the functional shading mode—so be sure to have it switched on before setting the new groups. Remember, \shadingmode{functional} without an optional parameter clears all groups defined before, see above. The following example shows the definitions needed to produce an output which is identical to the functional mode 'charge':

```
\begin{texshade}{\langle alignmentfile \rangle}
```

```
\shadingmode{functional}
 \funcgroup{acidic ($-$)}{DE}{White}{Red}{upper}{up}
 \funcgroup{basic ($+$)}{HKR}{White}{Blue}{upper}{up}
\end{texshade}
```

4.3 Appearance of the consensus line

An important parameter for the calculation of the consensus is the threshold percentage. Default setting is 50%, i.e. to become the consensus residue more than half of the residues at this position must be identical or similar, depending on the shading mode. Any percentage between 0 and 100 is allowed and can be set with $\threshold{percentage}$, e.g. $\threshold{50}$.

Additionally, an optional parameter can be set, e.g. \threshold[90] {50}, to label residues that are highly conserved in a special color (see example on page 15).

Another possibility is to set one sequence of the alignment as consensus and compare the other sequences to this one. Therefore, the command $\constosingleseq{\langle seqref \rangle}$ is provided. The $\langle seqref \rangle$ selects the sequence to be used as consensus (numbering according to the appearance in the alignment file; top sequence is number 1, or use the sequence name). Nevertheless, the threshold percentage is also taken into account, i. e. with a threshold of 50% half of the sequences must be identical or similar compared to the specified consensus sequence in order to be shaded. With \constoallseqs the consensus is calculated considering all sequences (the case described in the paragraph above).

Consensus lines are displayed either on the top or at the bottom of the alignment by calling

with $\langle scale \rangle$ Gray, BlueRed, RedBlue, GreenRed, RedGreen, ColdHot (recommended), HotCold, or T-Coffee 1.5.3, and $\langle position \rangle$ top or bottom.

The first color defines the foreground, i.e. the letters, the second color—if specified—defines the background. If a color scale is named the consensus will be shaded according to the level of sequence conservation (see section 4.3.1 on residue weight tables below). For an example output see page 17. You can find more information on color scales on page 71. The calculated consensus colors can be exported as a Pymol [8] script by \exportconsensus[$\langle filename \rangle$] { $\langle seqref \rangle$ }. If no $\langle filename \rangle$ is specified export.txt will be used. The generated file can be opened in Pymol in order to shade a 3D model of the sequence $\langle seqref \rangle$.

To hide the consensus use **\hideconsensus**. The consensus line is named 'consensus' in english texts, 'consenso' in spanish or 'Konsen-

sus' if the german.sty is used. With $nameconsensus{\langle name \rangle}$ any name can be set.

You can tell $T_E X_{shad}$ which symbols or letters to use in the consensus line for different matching qualities by

 $\det consensus \{\langle symbol1 \rangle\} \{\langle symbol2 \rangle\} \{\langle symbol3 \rangle\}.$

The following parameters are allowed for symobols 1–3:

- 1. $\langle symbol1 \rangle =$ no match symbol (if below threshold)
 - any character or letter
 - {} (empty braces) for blank space
- 2. $\langle symbol 2 \rangle = conserved symbol (if threshold is exceeded)$
 - upper (prints the consensus residue in uppercase)
 - lower (prints the consensus residue in lowercase)
 - any character or letter
 - {} (empty braces) for blank space

3. $\langle symbol \rangle =$ highly conserved symbol (if

\allmatchspecial is active)

• see $\langle symbol2 \rangle$

Example: \defconsensus{{}}{*}{upper} does not show non matching residues in the consensus line, marks conserved residues with '*', and displays the uppercase letter of the consensus residue at positions with high conservation.

Finally, the colors of the above defined symbols are adjustable by the command:

 $\consensus colors \{ \langle res. col. 1 \rangle \} \{ \langle shad. col. 1 \rangle \} \\ \{ \langle res. col. 2 \rangle \} \{ \langle shad. col. 2 \rangle \} \\ \{ \langle res. col. 3 \rangle \} \{ \langle shad. col. 3 \rangle \}$

The color definitions are in the same order as in the **\defconsensus** command:

- 1. $\langle res.col.1 \rangle =$ no match residue color (if below threshold) $\langle shad.col.1 \rangle =$ no match background color
- 2. $\langle res.col.2 \rangle = \text{conserved residue color (if threshold is exceeded)}$ $\langle shad.col.2 \rangle = \text{conserved background color}$
- 3. $\langle res. col. 3 \rangle =$ highly conserved residue color (if

\allmatchspecial is active)

 $\langle shad.col.3 \rangle =$ highly conserved background color

For colors which are not to be changed empty braces can be used. Example:

\consensuscolors{}{}{Blue}{White}{Red}{Green}

Non matching symbol colors are not changed, conserved residues are displayed blue on white and highly conserved residues appear as red symbols on a green background in the consensus line.

4.3.1 Residue weight tables

The degree of similarity between two amino acid residues is defined using so-called *residue weight tables*. The values usually range roughly from -10 to 10, with positive values denoting similarity and negative values dissimilarity. The most simple table sets pairs of identical residues to a value of 10 and all others to 0, i.e. the identity matrix. Several more matrices based on extensive protein alignments exist and can be used, e.g. PAM250 (Point Accepted Mutations), PAM100, or BLOSUM62 (BLOcks of amino acid SUbstitution Matrix); for details see respective sources and section 6.3. T_EX**shade** further contains a structural matrix where similarity is defined on simple comparisons of the sidechain properties with respect to volume and hydropathy.

For calculation of the consensus color shading or for bar graphs or color scales in the feature lines (4.6.3), a residue weight table can be selected by $weighttable{\langle table \rangle}$ with $\langle table \rangle$ being identity, structural, PAM250, PAM100, or BLOSUM62 (default is identity). Which matrix suits the analysis best needs to be decided case by case. Due to the all-positive values of the structural matrix (section 6.3) the similarity level appears usually very high; the identity matrix simply represents the number of identical residues at each position. The PAM and BLOSUM matrices provide more differentiated results.

One can change individual values or even define his own weight table using the command $\setweight{\langle res.1 \rangle}{\langle res.2 \rangle}{\langle value \rangle}$, e.g. $\setweight{E}{Q}{2}$ or $\setweight{K}{C}{-5}$. A full table, thus, needs 200 entries (20 * 20/2). A value for the gap penalty is set with $\gappenalty{\langle value \rangle}$.

4.4 Display of logos

4.4.1 Sequence logos

In a sequence logo [7], the information content $I(P_i)$ of each alignment position *i* is defined as

$$I(P_i) = \log_2 |\Sigma| + \sum P_{ij} \cdot \log_2 P_{ij}$$

with $|\Sigma|$ being the cardinality of the used alphabet, i.e. 4 for DNA and 20 for protein sequences, and P_{ij} being the frequency of residue *j* at this position. Each position is displayed as a stack of residue symbols whose heights represent their proportion of the information content (example on p.34).

The display of sequence logos can be either on the top or at the bottom of a nucleotide or protein alignment. Logos will be shown after the command: $\showsequencelogo[\langle colorset \rangle] \{\langle top/bottom \rangle\}$. If no optional $\langle colorset \rangle$ is selected the residues will be shaded as follows:

- Nucleotide sequences
 - G : Black
 - A : Green
 - T,U : Red
 - C : Blue
- Protein sequences (similar to rasmol)
 - D,E : Red
 - C,M : Yellow
 - K,R : Blue
 - S,T : Orange
 - F,Y : MidnightBlue

- N,Q: Cyan
 - G : LightGray
- L,V,I : Green
 - A : DarkGray
 - W : CarnationPink
 - H : CornflowerBlue
 - P : Apricot
 - B,Z : LightMagenta

Optional color sets correspond to the functional shading modes chemical, rasmol, hydropathy, structure, standard area, accessible area (see p.48). The \showsequencelogo command can be reversed by \hidesequencelogo.

Logo colors can be turned to 'Black' with the command $\clearlogocolors[\langle color \rangle$ } with the optional parameter not set. The optional parameter can be used to set all residue colors to $\langle color \rangle$, e.g. $\clearlogocolors[Blue]$. User specific logo color sets are defined by using $\logocolor{\langle residues \rangle}{\langle color \rangle}$, e.g. $\logocolor{DE}{Red} \logocolor{CM}{Yellow}$ etc.

It is common practice for protein sequence logos to correct amino acid frequencies to the background frequency in the alignment, which usually differs from the equal distribution of 5% for each residue. Frequency correction can be turned on by \dofrequencycorrection and off by \undofrequencycorrection.

The vertical extent of the logo can be changed by $\logostretch{\langle factor \rangle}$, e.g. $\logostretch{1.5}$. The width of the logo characters is dependent on the character width set for the alignment, see \charstretch on p.62.

Finally, the bit-scale can be turned off and on using hidelogoscaleand $showlogoscale[\langle color \rangle] \{\langle position \rangle\}$, respectively, with $\langle position \rangle$ left, right, or leftright and an optional $\langle color \rangle$. A name for the sequence logo can be set, which is displayed next to the scale by $namesequencelogo \{\langle name \rangle\}$.

4.4.2 Subfamily logos

Subfamily logos provide a novel tool to visualize subfamily-specific sequence deviations at alignment positions with a high information content in an intuitive way [14].

This is achieved by subtracting from the frequency of a residue within a pre-defined subset of sequences, i. e. a subfamily, the frequency of this residue in the remaining set of sequences. The difference is then weighted by the information content, see above section on sequence logos. An example is shown on p.35.

Subtraction of frequencies produces values from -1 to 1. Positive values correspond to residues which are characteristic for the subfamily (shown upright in the output), negative values to those that are typical for the remaining sequences (shown upside-down). Positions with an equal distribution of the residue result in a zero value.

Subfamily logos are displayed analogous to sequence logos by the command $\showsubfamilylogo[\langle colorset \rangle] \{\langle top/bottom \rangle\}\$ and hidden by \bidesubfamilylogo . To calculate a subfamily logo, it is further required to define a subfamily within the alignment by $\setsubfamily\{\langle seqrefs \rangle\}\$, e.g. $\setsubfamily\{1-10,20,AQP3\}$.

For coloring residues, display/stretching of the scales, and frequency correction the same commands as for sequence logos apply with two exceptions. First, subfamily logos contain negative values, which can be displayed $\shownegatives[\langle weak, medium, strong \rangle]$ or hidden \bidenegatives . Without the optional parameter negative residues will be tinted by 50%, i.e. medium. This greatly improves readability. Second, a name for the subfamily logo is set by $\mbox{namesubfamilylogo}[\langle neg.name \rangle] \{\langle pos.name \rangle\}$ with a required name for the positive part of the logo and an optional name for the negative part.

In order to better recognize relevant positions in the subfamily logo, a bit-value can be set above which the deviation is considered relevant by the command $\relevance{\langle bit-value \rangle}$. If this command is not given 2.321 bit is assumed for proteins, i.e. $\log_2 5$, and 1 bit for DNA, i.e. $\log_2 2$. Such positions will be labeled by $\showrelevance[\langle color \rangle] \{\langle symbol \rangle\}$, e.g. $\showrelevance[Blue] \{\nabla\}\}$. The symbol will be hidden with \hiderelevance .

4.5 Appearance of the sequence lines

4.5.1 Names, numbers and gaps

Many parameters that influence the appearance of the actual sequence lines can be changed for customization. Thus, the sequence names can be shown colored via $\langle color \rangle$ either left or right by with $\langle position \rangle$ set to left or right. The numbering can be displayed either left or right and even on both sides by

with $\langle position \rangle$ left, right or leftright. Both, names and numbering can be displayed on the same side. The colors can also be set with \namescolor{ $\langle color \rangle$ } and \numberingcolor{ $\langle color \rangle$ }, respectively. TEXshade uses the sequence names from the alignment input file. This can cause some problems during the TEX-run when special characters are present in those names! TEXshade does not accept the following characters in sequence names: \ { } @ spaces and the tilde. Those have to be replaced in the input file. The characters # and % can only be used with a leading backslash, e.g. \#. This must also be changed in the input file. All other special characters should be displayed properly.

Sequence names that are accepted by $T_EX_{shad}e$ can further be changed in the texshade environment:

 $\langle seqref \rangle$ selects the sequence whose name is to be changed. The basis for the $\langle seqref \rangle$ is the appearance in the alignment input file with the top sequence = 1, or the old name. In order to change the colors only of some sequence names or numbers the commands $\mbox{namecolor}\{\langle seq1 \rangle, \ldots, \langle seqn \rangle\}\{\langle color \rangle\}$ and $\mbox{numbercolor}\{\langle seq1 \rangle, \ldots, \langle seqn \rangle\}\{\langle color \rangle\}$ are provided.

In order to hide all names or the numbering use the command **\hidenames** or **\hidenumbering**. If only the names or numbers of some sequences should be hidden apply

 $\lambda \in \{\langle seq1 \rangle, \ldots, \langle seqn \rangle\}$ or

 $\label{eq:linear} \$, ..., (seq n)}, respectively.

In some situations, e.g. when only sections of sequences are displayed, one may not want to have the residue numbering start 1. The out with number command $startnumber[\langle start..stop \rangle] \{\langle seqref \rangle\} \{\langle startnumber \rangle\}$ allows one to set the starting number of any sequence to any value incl. negative values but except '0' which is not usually used in sequence numbering (the transition from negative to positive values is like this: $\dots -2, -1, 1, 2 \dots$). If, however, the use of the number '0' is wanted as sometimes in sequence logos this can be turned on by \allowzero and off with \disallowzero. The optional parameter can be used to

truncate the sequence display to a certain section (see also **\setends** below).

TEX**shade** needs to know the correct length of the sequences to be able to break them right after the last residue. If MSF files are used as an input the length is already given but the calculation is usually wrong because the gaps are also counted. Thus, TEX**shade** counts the number of residues during each run by itself and stores the values in the .aux file. That means that it needs two runs to get the numbers right. Again, this is only important if the gap symbol after the sequence end should be suppressed, see below (hideleadinggaps).

If you know the correct length of the sequences you can use the command

in order to set the values by hand and have the gaps break properly already in the first $T_{\rm E}X$ run.

Example: $\ensuremath{seqlength{1}{346}}\ensuremath{means}\xspace$ that sequence no. 1 is 346 residues long.

Another possibility to label sequence positions is to switch on a ruler on the top or at the bottom of the sequence block using $\showruler[\langle color \rangle] \{\langle position \rangle\} \{\langle seqref \rangle\}$. The residue ruler of one sequence $\langle seqref \rangle$ or the consensus (declare 'consensus' as $\langle seqref \rangle$) can be displayed at $\langle position \rangle$ top or bottom. The ruler is hidden with \hideruler . The steps between two numbers are set by $\rulersteps\{\langle number \rangle\}$. If the steps are set to be very close (< 4) or when every position is numbered, the numbering is automatically rotated by 90°. Using \rotateruler and \unrotateruler this can be done and undone manually. In order to change the ruler color use the optional parameter or the command $\rulercolor\{\langle color \rangle\}$. Also, the label and its color at individual ruler positions can be changed by the user to a string using $\neuerulerpos\{\langle number \rangle\}\{\langle text \rangle[\langle color \rangle]\}$ (see example on p. 24).

Further, the symbol which is displayed in sequence gaps is freely selectable with \gapchar{(symbol)}. (symbol) can be any character or symbol. If math symbols are to be used math mode must be activated by \$ characters, i.e. \gapchar{{\$\triangle\$}}. Note the double curly braces in the last command. Everytime a 'complex' character is used, i.e. a character definition consisting of more than one letter, it must be braced in order to be interpreted as one character. One exception is \gapchar{rule}; with this parameter lines are drawn in the sequence gaps with a certain thickness defined by $gaprule{\langle thickness \rangle}$, e.g. $gaprule{1.5pt}$. The colors of the gaps and gap symbols are set by $gapcolors{\langle symbol color \rangle}{\langle background color \rangle}$.

There are some discussions whether or not to display gap symbols before and after the actual sequence. Since v1.3a one can control the appearance of those gap symbols by the commands **\showleadinggaps** and **\hideleadinggaps**. By default, leading gaps are indicated by symbols despite my personal thinking that it could suggest that there are some not displayed residues upstream resp. downstream of the gap.

At certain instances a protein alignment input file may contain stop positions, e. g. due to frame shifts in the underlying DNA sequence. If such positions are labeled in the input with an * this will be shown in the output as well as an asterisk, i.e. distinguishable from a normal gap symbol. The character shown at stop positions can be changed by $stopchar{\langle symbol \rangle}$.

4.5.2 Displaying selected residues in the alignment

 T_EX **shade** can display a section of the complete alignment without the need to edit the alignment input file or even to re-calculate the entire alignment. This allows one to use one single alignment of the full length proteins or open reading frames for multiple visualizations of different sections in a document as done in this manual. Thus, the file AQPpro.MSF contains the full-length multiple protein alignment of five aquaporins but only sections are displayed as examples in 1.5.1 through 1.5.5. The definition of a section is done by

 $\ [\langle startnumber \rangle] \{\langle start..stop \rangle\}.$

Again, $\langle seqref \rangle$ is the sequence number based on the appearance in the alignment file, or the name; further, in order to use the consensus as a measure for the sequence section the string 'consensus' as $\langle seqref \rangle$ is accepted. The specified sequence is truncated at positions $\langle start \rangle$ and $\langle stop \rangle$. All other sequences are cut accordingly. If the number of the first residue in the sequence is set to a new value with the \startnumber command (s. a.) this is taken into account. The $\langle startnumber \rangle$ can be set as an optional parameter directly in the \setends command as well.

Some examples:

a) \setends{1}{21..100}

b) \startnumber{1}{101} \setends{1}{121..200}

Both commands select the same sequence section from the alignment but numbering for sequence 1 starts at position 21 in the first example and at position 121 in the latter.

- c) \setends[101]{1}{121..200} equals example b.
- d) $startnumber[121..200]{1}{101}$ also equals example b.

```
e) \setends{consensus}{21..100}
```

This may describe a very different section of the multiple sequence alignment because the consensus counts every position including gaps. The output can be even further restricted to individually selected residues, e.g. to eliminate uninteresting alignment stretches or to condense the output, by:

Here, $\langle seqref \rangle$ denotes the reference sequence by its number or name. This sequence is used to define the alignment positions $\langle selection \rangle$ to be shown. $\langle selection \rangle$ can have two different formats depending on whether (a) the user wants to select the positions manually or (b) T_EX**shade** is supposed to select the residues based on 3D coordinates provided by a PDB file.

To select the residues manually, the user provides a position list of the following format:

 $\{\langle start1 \rangle .. \langle stop1 \rangle, \langle start2 \rangle .. \langle stop2 \rangle, ..., \langle start n \rangle .. \langle stop n \rangle \}$

For how to select positions by 3D coordinates provided by a PDB file, see 4.6.1.

Examples (see also p. 15ff):

```
\setdomain{1}{20..80}
\setdomain{consensus}{20..80,100..150,200..220}
\setdomain{AQP1}{point[6]:1FX8.pdb,173[side]}
\setdomain{3}{plane[0.5]:1JN4.pdb,66[CA],73[side],199[CA]}
```

It is helpful to show a ruler (probably single-stepped, see p. 58) to label the residue positions.

The resulting gaps between sequence stretches are marked by a vertical rule, which can be changed in thickness by

 $\operatorname{domaingaprule}(\operatorname{thickness}), e.g. \operatorname{domaingaprule}(1pt).$

Also, the colors can be set by

$\operatorname{domaingapcolors}{\langle foreground \rangle}{\langle background \rangle}$

e.g. \domaingapcolors{Blue}{Yellow}.

4.5.3 Hiding, killing, separating and ordering

If one or more sequences from the alignment input file should be used for the calculation of the consensus but it is desired not to display these sequences in the final output use the command $\frac{\frac{\sqrt{seq1}}{\sqrt{seq2}}, \dots, \frac{\sqrt{seq n}}{}$. For consecutive sequence numbers a dash can be used, e.g. $\frac{1-3}{\frac{1}{2}}$ instead of $\frac{1-3}{1}$. Decending series are also permitted, e.g. $\frac{1-3}{1}$. This command allows one for example to hide the sequence which has been defined as the consensus sequence with $\frac{\sqrt{seq2}}{\sqrt{seq2}}$. When all sequences should be hidden, e.g. to show a sequence logo alone, one can simply say $\frac{1}{\sqrt{seq2}}$. This command is reversed by $\frac{\sqrt{showseqs}}{\sqrt{showseqs}}$.

In order to completely exclude sequences the command $\killseq{\langle seq1 \rangle, \langle seq2 \rangle, ..., \langle seq n \rangle}$ is provided. Again, for number series the dash can be used (s. a.). The designated sequences are neither displayed nor considered for the calculation of the consensus. This is another possibility to comment out sequences in addition to the use of an exclamation point infront of the Name: definition in an MSF-file (see figure on page 40).

The command $\donotshade{\langle seq1 \rangle, \langle seq2 \rangle..., \langle seqn \rangle}$ makes one or more sequences (remember the dash, s. a.) appear unshaded in black letters on white background. This does not influence any other sequences or the consensus calculation.

If a very graphical output of the sequences is desired, the residue symbols or letters can be blanked out by **\hideresidues**. Now, only the shaded boxes are printed. In combination with **\gapchar{rule}** one obtains alignments in a style à la Mondrian. The residues reappear with **\showresidues**.

If an alignment contains members of several subgroups of a protein or a gene family it may be rather helpful to visualize the group divisions by a separation line. Therefore, the command $\separationline{\langle seqref \rangle}$ is applicable. This command inserts vertical space after the sequence which is referred to by $\langle seqref \rangle$. How much space is inserted is defined by one of the following commands: \smallsep , \medsep (default) or \bigsep . These lengths correspond to the known $\mbox{small-}, \mbox{med-} and \mbox{bigskip} commands. With <math>\mbox{vsepspace}{\langle length \rangle}$ any length with any T_EX unit can be assigned, e.g. $\mbox{vsepspace}{2mm}$.

The sequence order given by the alignment input file is easily reorganized by $\orderseqs{\langle seq1 \rangle, \langle seq2 \rangle, ..., \langle seqn \rangle}$ without the need for editing the alignment input file (which would be a big copy'n'paste job). Make sure that all sequences are assigned in this command. If there are more sequences present than numbers or names in the command an error message will occur. Here also, the dash can be used for sequence number series. Example: $\orderseqs{1-3,6-4,7}$ is equivalent to $\orderseqs{1,2,3,6,5,4,7}$. Reordering of sequences only changes the output; all commands using the parameter $\langle seqref \rangle$ are not influenced, because $\langle seqref \rangle$ always corresponds to the appearance in the alignment file. Thus, to completely reverse the order of a five sequence alignment simply type $\orderseqs{5-1}$.

4.5.4 Residues per line and further settings

By default T_EX shade puts the highest possible by five divisible number of residues in one line depending on the \linewidth . With $\residuesperline{\langle number \rangle}$ a new value can be set. If this value exceeds the highest possible number of residues per line it is ignored; lower values are accepted of course. But also in the latter case the number of residues printed per line is rounded such to be divisible by five. To force T_EX shade to set lines with exactly the desired number of residues use the asterisk-extended command $\residuesperline*{\langle number \rangle}$. You have to take care yourself of the alignment width after this command, because in this mode T_EX shade does not check the length of the lines any more.

T_EX**shade** calculates the dimensions of a shaded box from the width and height of the uppercase letter 'M' and the depth of the lowercase 'g'. Depending on the font used for the sequence residues the box dimensions might not be fully satisfactory. With $\charstretch{depth} depth$, respectively, of the boxes can be multiplied individually by a $\langle factor \rangle$ to stretch (> 1) or shrink (< 1) the dimensions.

The reserved space for the sequence numbering is set by the command $\mberingwidth{\langle n \ digits \rangle}$. Here, the default setting is four-digit numbering, i.e. -999 through 9999. If this range is to be changed assign the desired number as parameter $\langle n \ digits \rangle$, e.g. $\mberingwidth{111111}$ reserves space for 6 digit numbering. The vertical space between the sequence blocks can be controlled by the commands $\mbox{smallblockskip}$, $\mbox{medblockskip}$ (default setting), \bigblockskip or $\mbox{noblockskip}$. Further, the command $\bigblockspace{\langle length \rangle}$ allows one to set a defined space length using any TEX unit, e.g. $\blockspace{0.4in}$.

Two more commands set the space between the sequence blocks to be flexible (\flexblockspace) (default) or fixed (\fixblockspace). Flexible means, that only the vertical white space between the blocks is kept to the settings by e.g. \medblockskip. This results in flexible space between the actual blocks depending on the presence of feature lines. When switching to fixed space the distance of the blocks is kept constant by using more white space between blocks without feature lines. Thus, a difference between flexible and fixed space will only be noticeable when features are used.

The position of the output can be aligned left, right or centered on the page by $\label{eq:position}$ with the $\langle position \rangle$ parameter left, center or right.

4.5.5 Fingerprinting

An easy way to gain an overview on complete alignments is provided by displaying a so called alignment 'fingerprint'. In this style the whole sequence can be shown in one line. Due to the lacking space the residue names are hidden and the shaded boxes are reduced to thin vertical colored lines. The command $fingerprint{res. per line}}$ takes one argument stating the desired number of residues per line, e.g. fingerprint{1000}. All TEX**shade** commands are compatible with fingerprint, i.e. all shading modes are applicable for displaying overviews on similarity or every functional aspect. Also, all kinds of labeling—as described in the following—work with this command.

4.6 Individual shading and labeling of sequence stretches

Computer calculated conservation shading is informative—but even more information can be visualized by additional labeling of positions and regions of interest with different colors, text styles or graphical marks and descriptive text. All this is provided by easy to handle $T_{\rm EX}$ shade commands.

4.6.1 Shading of regions and blocks

Besides the shading calculated by T_EX shade any region can be additionally shaded with a color specified by the user. This is very useful to highlight secondary protein modification sites such as phosphorylation or glycosylation sites, or longer motifs for example protein/protein interaction sites or protein domains. This is done with the following command:

Here, $\langle seqref \rangle$ refers to the sequence by its name or number within the alignment. The foreground and background colors can be set with the last two parameters. $\langle selection \rangle$ can have three different formats depending on whether (a) the user wants to select the positions manually, (b) the selection should be based on a sequence motif to be found by T_EX**shade**, or (c) T_EX**shade** is supposed to select the residues based on 3D coordinates provided by a PDB file.

(a) To select the residues manually, the user provides a position list of the following format:

```
\{\langle start1 \rangle .. \langle stop1 \rangle, \langle start2 \rangle .. \langle stop2 \rangle, ..., \langle start n \rangle .. \langle stop n \rangle \}
```

In order to shade residue number 13 and the region 20–30 of sequence number 1 in red letters on a green background type the following command:

```
\shaderegion{1}{13..13,20..30}{Red}{Green}
```

If the consensus is to be shaded use consensus as $\langle seqref \rangle$.

(b) A sequence motif can be given, which will be found and labeled. A simple example selecting all sequence motifs 'NNAD' in sequence '1' would be:

\shaderegion{1}{NNAD}{Red}{Green}

The definition can further include 'X' for any residue and groups of residues in brackets at uncertain positions. The example below will find all motifs in sequence '1' that start with an asparagine, followed by any two residues, an acidic, a basic residue, again any two residues and finally a glutamine.

```
\shaderegion{1}{NXX[DE][KR]XXQ}{Red}{Green}
```

(c) In order to select positions based on the 3D structure a PDB structure file is required. T_FX**shade** can select residues within a given

distance in Å around a point, along a line, or above and below a plane, which are defined by one to three residues. The points can be further specified to be the α -carbon atom, i.e. the protein backbone, or the most distant atom of the sidechain. Accordingly, $\langle selection \rangle$ has one of the formats:

```
 \{point [\langle dist \rangle] : \langle file \rangle, \langle num \rangle [CA/side] \} \\ \{line [\langle dist \rangle] : \langle file \rangle, \langle num1 \rangle [CA/side], \langle num2 \rangle [CA/side] \} \\ \{plane [\langle dist \rangle] : \langle file \rangle, \langle num1 \rangle [CA/side], \langle num2 \rangle [CA/side], \langle num3 \rangle [CA/side] \} \\ \} \\ \{num3 \rangle [CA/side] \} \\ \{num3 \rangle [CA/side] \} \\ \} \\ \{num3 \rangle [CA/side] \} \\ \} \\ \{num3 \rangle [CA/side] \} \\ \{num3 \rangle [CA/side] \} \\ \} \\ \{num3 \rangle [CA/side] \} \\ \{num3
```

Example: in order to select and shade as above all residues that are within an 8 Å sphere around the α -carbon of residue 81 and the data are provided in the PDB file 1J4N.pdb, type:

```
\ \
```

Example: two points denote a line, hence, give two residues to select everything within 1 Å along the line between the α -carbon of residue 81 and the sidechain of residue 168 with:

```
\shaderegion{1}{line[1]:1J4N.pdb,81[CA],168[side]}
```

{Red}{Green}

Definition of a plane follows the same format but requires three points. If the optional parameters [$meta{dist}$] and [CA/side] are not given, T_EX shade assumes [1] and [side], respectively.

In case one needs the position numbers of the selected residues for usage in other applications, those can be either printed in the T_EX document with $\printPDBlist{\langle selection \rangle}$ or shown during the T_EX run with $\mbox{messagePDBlist}(\selection \rangle)$. The commands can be used outside of the T_EX shade environment.

Both selection formats, i.e. a manually given list and the 3D selection, can be used with \block (see below), \tintregion , \tintblock , \emphregion , \emphblock , \lowerregion , \lowerblock , \frameblock (all in 4.6.2), and \feature (4.6.3).

In analogy to \shaderegion which is restricted to a single sequence, \shadeblock shades the corresponding region in all other sequences as well except the consensus. If also the consensus is to be shaded define the region using consensus as $\langle seqref \rangle$.

Another option is to change the whole set of shading colors for certain sequence blocks, e. g. from **blues** to **reds** or self-defined color sets (see 4.1). Therefore, the following command was implemented:

Examples:

```
\changeshadingcolors{1}{10..50}{reds}
```

```
\changeshadingcolors{AQP1}{[AS]NKD}{my_set}
```

etc.

4.6.2 Emphasizing, tinting, lowercasing, and framing

If it is preferred to keep the calculated shading colors but distinct regions or blocks are yet to be emphasized one can use the following commands to change the font style of such regions:

 $\ensuremath{\mathsf{emphregion}}{\langle seqref \rangle} \{\langle selection \rangle\}$

and

```
\mathbb{C} \left( seqref \right)
```

For the format possibilities of the $\langle selection \rangle$ parameter please see 4.6.1.

```
Which style T_EX and e uses for emphasizing regions is defined by 
\emphdefault{\langle style \rangle}. Default setting is the italics font shape (set by \emphdefault{it}). In order to change this setting choose one of the styles bf, md, up, it, sl, rm, sf, tt. Example: \emphdefault{bf}
```

Further, it is possible to tint the region or block in question or to switch the characters to lowercase by using the commands (for example see hydropathy-figure on page 21):

```
\tintregion{\langle seqref \rangle}{\langle selection \rangle}
```

and

```
\tilde{\langle seqref \rangle} \{\langle selection \rangle\}
```

The level of tinting in the region in question can be set by $tintdefault{\langle level \rangle}$ with weak, normal, and strong as possible $\langle level \rangle$ s.

and

 $\lowerblock{\langle seqref \rangle}{\langle selection \rangle}$

Another option is to draw a bounding box around the sequence block in question (for an example see diversity mode-figure on page 20) with the command:²⁴

 $frameblock{\langle seqref \rangle}{\langle selection \rangle}{\langle color \rangle}{\langle length \rangle}}$

With the optional parameter the default line thickness of the frame can be changed, example: $frameblock{1}{10..20,50..70}{Red[2pt]}$

4.6.3 Graphical labeling of sequence features

The **\feature** command is designed to fulfill most needs for the graphical labeling of sequence stretches and the setting of descriptive text. It needs five parameters:

 $feature{\langle position \rangle}{\langle seqref \rangle}{\langle selection \rangle}{\langle labelstyle \rangle}{\langle text \rangle}$

In the following paragraphs all possible parameter settings of this rather complex but mighty command are discussed in detail. The parameter $\langle position \rangle$ tells T_FXshade where to display the feature label, i.e. on the top of the alignment (top), or at the bottom (bottom). Further, there can be three more feature lines ontop of the top feature line (ttop, tttop, and ttttop) or below the bottom feature line (bbottom, bbbottom, bbbbottom). Thus, up to eight features overlapping in eight different lines may be displayed. Depending on the content of the feature lines the gaps between them might be not satisfactory. Therefore, eight separate commands can be employed to change the space below ttttop, tttop, ttop, or top $(\topspace{\langle length \rangle} etc.)$, and above bottom, bbottom, bbottom, or bbbbottom (\bottomspace{(length)} etc.). Use positive values to further separate the lines, e.g. \ttopspace{3mm} or negative values to reduce the space, e.g. \bottomspace{-0.1in}.

The argument $\langle seqref \rangle$ and the third parameter containing the $\langle selection \rangle$ of the specified residues are identical to the ones described before in several commands, e.g. \shaderegion (4.6.1).

New is the fourth parameter for the definition of the label style. There are many possibilities like braces, helices, boxes, arrows, bars, any fill character, bar graphs, color scales or even translations of the specified regions.

Braces:

In order to display an over- or underbrace as a label use the parameter

²⁴Thanks to Alan Robinson for inspiration.

{brace}. Depending on the $\langle position \rangle$ (ttttop, tttop, ttop, top, bottom, bbottom, or bbbbottom) the respective brace is displayed. The standard color of braces is black. It can be changed by an optional parameter directly after the definition of the symbol, e.g. {brace[Red]}.

Protein α -Helices:

The parameter $\{\text{helix}\}$ will plot a symbolized α -helix as a label. The standard color of the helix spiral is black. It can be changed by an optional parameter directly after the definition of the symbol, e.g. $\{\text{helix}[\text{Red}]\}$.

Filling a stretch with a symbol:

A region can be filled with any character for labeling purposes using the parameter $\{fill: \langle symbol \rangle\}$. The $\langle symbol \rangle$ is freely selectable; the usage is like in $\langle gapchar (4.5.1)$. Do not use spaces before or after the expression $\langle symbol \rangle$; this will shift the symbols to the respective direction. The standard color of the fill symbol is black. It can be changed by an optional parameter directly after the definition of the symbol, e.g. $\{fill:\Let\[Red]\}$.

The **\feature** command does not like special characters in text mode, e. g. **\dag**. One has to use the math version of those symbols between **\$**-signs. The following quite common text symbols have also a math equivalent²⁵:

symbol	command	description
†	\$\dagger\$	dagger
‡	\$\ddagger\$	double dagger
¶	<pre>\$\mathparagraph\$</pre>	paragraph mark
§	\$\mathsection\$	section mark
\$	\$\mathdollar\$	dollar
{	<pre>\$\lbrace\$</pre>	left brace
}	\$\rbrace\$	right brace

Labeling restriction or protease cutting sites:

If a label is needed that points between two residues, e.g. for showing restriction sites, simply use the feature style {restriction[$\langle color \rangle$]}. This will show a filled triangle with the tip right between the residues to be labeled, e.g. \feature{top}{1}{25..26}{restriction[Blue]}{EcoR I}.

²⁵Thanks to Darrell Conklin for reporting this problem

Boxes:

Boxed text is printed using the parameter $\{box: \langle text \rangle\}$. By default black letters in a white framed box are displayed. In order to change these colors optional parameters can be included in the argument:

 $box[\langle framecolor, boxcolor \rangle][\langle length \rangle]: \langle text \rangle[\langle textcolor \rangle]\}.$

If the box frame and fill colors are the same it is sufficient to use only this one color as an argument in the command. The optional parameter $\langle length \rangle$ defines the thickness of the box frame. If this parameter is not set in the command the value from the $featurerule{\langle length \rangle}$ command (see below) is used.

Examples:

{box[Blue]:\$\alpha\$-helix[Yellow]}
{box[Blue,Red]:\$\alpha\$-helix[Yellow]}
{box[Blue,Red][2pt]:\$\alpha\$-helix[Yellow]}

Horizontal bars and arrows:

For displaying bars and arrows a simple selection scheme consisting of three consecutive characters is used as the $\langle labelstyle \rangle$ parameter. Each bar or arrow is defined by its left end, the middle part, and the right end. The following table gives some examples for the construction of arrows and bars.

middle	
$\text{left end} \downarrow \text{right end}$	
	plain bar
===	double bar
>	right arrow
'->	right arrow with up hook
<-	left maps to arrow
<-0	left arrow with ball at right end
<=>	double arrow, two heads
, — ,	plain bar with down hooks
=	double bar with vertical ends

All combinations of the left-end-characters (-=<', $|o\rangle$, the middlecharacters (-=), and the right-end-characters (-=>', $|o\rangle$) are allowed and produce the desired arrow or bar. The color is changed as described above. The thickness can be generally set by the separate command \featurerule{ $\langle length \rangle$ } with any TEX measure as $\langle length \rangle$, e.g. \featurerule{3pt}. This value is then used for all arrows, bars, and boxes (see above) throughout the alignment. If an individual thickness for a particular arrow should be set one can add an optional parameter to the *\labelstyle* parameter, e.g. {o->[Red][1mm]}. Similar to the boxes described above, a text can be put on the arrow or bar, e.g. {<->[Red][1mm]:\$\beta\$-sheet[Blue]}.

In T_EX**shade** versions before v1.9, the original $\text{LAT}_{E}X$ -arrows were used. These have now been replaced by more modern looking arrows with scalable line thickness. If the classical look is requested, use v instead of < or > in the arrow definition, e.g. $\{--v\}$, to get them back. The new arrow style makes use of of the AMS math symbol font (amssymb.sty). Thus, in order to display the arrow heads correctly make sure that this style is present on your system (usually it is in a common LAT_EX installation).

Sequence translations:

With the option {translate}, sequence stretches can be translated from nucleotide to peptide sequences as well as backtranslations from peptide to nucleotide sequences are possible. Default setting for the translations is the standard genetic code. Of course, the codons can be re-defined by the user. The command $\codon{(amino$ $acid)}{(triplet1, ..., triplet n)}$ has been implemented for this issue. The usage is simple. Replace (amino acid) by the single letter code of the amino acid to be defined and add a list of triplets for this residue. Example definition for the amino acid *alanine*:

\codon{A}{GCA,GCG,GCC,GCT,GCU,GCN}

Note the last triplet in the list. It contains an ambiguity code N which stands for *any* nucleotide. This triplet has been added at the last position because the last triplet is used for the generation of the backtranslated nucleotide sequence from a peptide. Two files are included in the TEXshade distribution as examples (standard.cod, ciliate.cod). If you want to define a new genetic code store your commands in a file like the examples. Such files with the suffix .cod can be loaded in the TEXshade environment by \geneticcode{ $\langle filename \rangle$ }, e.g. \geneticcode{ciliate}. Do not designate the suffix .cod in $\langle filename \rangle$. Please note, when inspecting the example files, that only the exchanges compared to the standard code need to be defined in a new genetic code file.

When DNA sequences are translated to protein the resulting amino acids are aligned to the second nucleotide of each triplet. It is more difficult to produce a satisfactory display of backtranslated nucleotide sequences due to the lack of space. You need thrice as much space than the original peptide sequence, because single letter amino acid code is translated to a triplet code. Therefore, the user can choose from five display styles for backtranslations depending on personal preferences:

 $backtranslabel[\langle size \rangle] \{\langle style \rangle\}, with$

$$\{\langle style \rangle\} = \{$$
horizontal $\}$
= {alternating}
= {zigzag}
= {oblique}
= {vertical}

 $\langle size \rangle$ can be any T_EX size from tiny up to Huge, but tiny is recommended (and default setting). Translations can be colored as all other labels, see above.

Bar graphs and color scales:

Sequence related numeral data, such as hydropathy or solvent accessibility data etc., can be shown in a feature line as bar graphs or color scales. The data are (a) pre-defined or calculated by T_EX shade due to amino acid properties or conservation, (b) are provided in a separate file or (c) may be entered by hand in the \feature command.

(a) Currently, three different properties can be plotted, i.e. hydrophobicity, molweight, and charge. Further, the level of sequence conservation at the given protein sequence stretch can be shown (conservation). See 4.3.1 for selecting an appropriate *residue* weight table.

(b) The format of a data file is simple: every value must appear in a separate line. Numbers and the Java-typical 'NaN' for 'Not a Number' are permitted. Comments are allowed, because $T_EXshade$ ignores all lines starting with a letter except 'NaN' lines (avoid '-' as the first character of a comment line as this is interpreted as a negative number). Make sure that there are as many values as positions defined as the sequence stretch in the feature command. $T_EXshade$ will read this file and determine the minimal and maximal values. These data are then normalized for plotting. Due to T_EX 's limited calculation capabilities no values above 10737 are allowed and the difference between minimum and maximum must not exceed this very number. Values below 0.001 may be susceptible to major rounding errors. Thus, try to

provide your data already normalized to moderate scales, e.g. 0.0-1.0 or -100-100.

(c) Data which is directly entered in the **\feature** command must be normalized to integer values with a maximal difference of 100 between the highest and lowest value, e.g. -50-50 or 0-100.

For (b) and (c), the range to be plotted can be set by hand as an This can be necoptional parameter in the \feature command. essary when the data file contains values between e.g. -0.44 and 0.87. Without help T_{FX} shade will assume -0.44 as minimum and 0.87 as maximum. But if the actual range to be plotted should be -1.0-1.0 this needs to be set manually, see examples below. Be aware of the fact, that if you define a scale by hand, which is more narrow than the values of the input, this will stretch the bars accordingly. It is NOT recommended to use this method for stretching bars vertically. Instead another command has been introduced. The plotted bars can be stretched by a factor if the appearance is not as desired: $\bargraphstretch{\langle factor \rangle}$. Here, the factor is multiplied with the bar length, e.g \bargraphstretch{2} will double the bar height, \bargraphstretch{0.5} will make them half as high. Similarly, color scales can be stretched vertically with $\colorscalestretch{\langle factor \rangle}.$

The default color of bar graphs is gray and can be changed by an optional parameter at the end of the label definition. Further, an optional background color can be chosen for the bars. Doing so will visualize the maximal bar extension.

Default for color scales is a 5% gray scale from very light gray to black (Gray). More colorful scales have been implemented, i.e. BlueRed, RedBlue, GreenRed, RedGreen, ColdHot and HotCold, the latter two being particularly useful for ranges from negative to positive values. Further, a scale called T-Coffee is available if T-Coffee shading information has been imported as the \shadingmode 4.1.

The general format of this feature label definition for bar graphs is:

 $bar[\langle min \rangle, \langle max \rangle]: \langle properties/file/data \rangle [\langle color(, bgcolor) \rangle]$

and for color scales:

 $\{color[\langle min \rangle, \langle max \rangle]: \langle properties/file/data \rangle[\langle scale \rangle] \}$

Some examples:

{bar:conservation}

{bar:conservation[T-Coffee]}

```
{bar:hydrophobicity}
{bar:charge[Red]}
{bar:molweight[Red,Gray10]}
{bar:10,20,30,40,50[Red]}
{bar[-20,40]:-10,0,10,20,30[Red,Gray10]}
{bar:data.txt}
{bar[-10,10]:data.txt[Red,Gray10]}
{color:conservation[BlueRed]}
{color:charge}
{color:hydrophobicity[GreenRed]}
{color:charge}
{color[-10,10]:data.txt[ColdHot]}
{color[-0.1,0.1]:otherdata.txt[ColdHot]}
```

See also the example output in section 1.6 on page 27.

No graphical label, only text:

If no graphical label is wanted the fourth parameter of **\feature** can be empty braces.

Finally, the fifth parameter of the **\feature** command contains the descriptive text for the labeled region. Type whatever you want incl. symbols and math chars. The text field can also contain sequence translations. In this case just set $\langle text \rangle = \{\texttt{translate}\}$. There is a command for setting the size and style of backtranslated sequences in the feature $\langle text \rangle$ which corresponds to the one described above:

```
backtranstext[\langle size \rangle] \{\langle style \rangle\}
```

Again, the color can be set by an optional parameter appended to the text. For how to change the font size of text or symbols in the feature style line (featurestyles) or the in descriptive text line (features) see section 4.9.1, page 79.

Another set of commands can be used to set a name for a feature line, which is printed together with the sequence names at the left or right side of the alignment, i.e.

 $\blue \label{eq:linear} $$ \blue \end{tilter} $$ \blue \end{tilt$

Using \showfeaturename will print the name in the same line as the descriptive text of the feature whereas \showfeaturestylename will put the name in the same line as the feature symbols. The color of such names can be generally changed with

```
featurenamescolor{(color)} and
```

 $featurestylenamescolor{(color)}$

or individually with

 $featurenamecolor{\langle ttttop...bbbbottom \rangle}{\langle color \rangle}$ and

 $featurestylenamecolor{\langle ttttop...bbbbottom \rangle}{\langle color \rangle}$

See section 5 for how to select colors in T_EX shade. Font styles can be set as usual (see section 4.9.1), e.g.

\setsize{featurenames}{large} or \featurestylenamesrm etc.

Examples for the appearance of features are given in the overview section (1), see:

similarity mode (1.5.2): fill-character; here, only one position is labeled. It is also possible to label a longer stretch, then, the character is printed several times to fill the specified region.

```
\feature{top}{1}{93..93}{fill:$\downarrow$}{first...}
\feature{bottom}{1}{98..98}{fill:$\uparrow$}{second...}
```

T-Coffee mode (1.5.3): T-Coffee color scale

```
\feature{top}{1}{30..63}{color:conservation[T-Coffee]}{}
\showfeaturestylename{bottom}{cons}
```

diversity mode (1.5.4): frames, text only
 \feature{top}{1}{77..109}{}{AQP2 species variants}
 \frameblock{1}{82..82,106..106}{Red[1pt]}

functional mode (1.5.5): bar graph, color scale, tinting, box, arrow, translation, brace, helix

```
\feature{top}{3}{153..165}
{bar[-50,50]:-50,-45,-40,...,40,45,50}{}
\feature{top}{3}{167..186}
```

```
{color:5,10,15,...,90,95,100[ColdHot]}{}
\feature{top}{1}{158..163}{brace}{tinted}
\tintblock{1}{158..163}
\feature{top}{1}{138..157}
        {box[Blue,Red][0.5pt]:$\alpha$-helix[Yellow]}
        {transmembrane domain 4}
\feature{top}{1}{164..170}{o->[Red]}{trans. dom. 5}
\feature{top}{1}{158..163}{translate[Blue]}{}
\backtranslabel{oblique}
\feature{bottom}{1}{158..163}
        {brace[Blue]}{loop D[Blue]}
\feature{top}{1}{158..163}{---}{loop}
\feature{tom}{}
```

bar graphs and color scales (1.6): sequence conservation, charge, molecular weight, hydrophobicity

4.6.4 Including secondary structure information

The DSSP [9], STRIDE [10], PHD [11] and HMMTOP [12] algorithms produce secondary protein structure predictions. PHD files contain both, secondary structure information and topology data. This information can be displayed in an alignment by one of the commands:

\includeDSSP	sec. structure calculated by DSSP
\includeSTRIDE	sec. structure calculated by STRIDE
\includePHDsec	sec. structure calculated by PHD
\includePHDtopo	topology data calculated by PHD
\includeHMMTOP	topology data calculated by HMMTOP

The syntax is $\includeDSSP{\langle seqref \rangle}{\langle filename \rangle}$, with seqref indicating the number or name of the sequence for which the secondary

structure data is calculated and **filename** designating the corresponding structure file to be included.

Several types of secondary structures are predicted by these programs; in order to designate them in T_EX **shade** use the names from the right column:

secondary structure	designation
DSSP and STRIDE	
4-helix (α -helix) isolated β -bridge extended strand (β -strand) 3-helix (3_{10} -helix) 5-helix (π -helix) H-bonded turn	alpha bridge beta 3-10 pi turn
PHDsec	
helix sheet	alpha beta
PHDtopo and HMMTOP	
internal region external region transmembrane domain	internal external TM

By default all three types of helices and the strands are displayed whereas turns and bridges are skipped. If it is desired to shown them as well, call for example \shownonDSSP{bridge,turn}. In analogy to this example all structure features can be activated in DSSP, STRIDE, PHDsec, PHDtopo and HMMTOP. In order to hide certain structure types use for example \hideonDSSP{3-10,pi}.

The DSSP format has two columns of sequence numberings. The first column is consecutive, whereas the second column contains the actual sequence numbering. This can be different from the first column when sequence parts are missing in the DSSP file. One can choose which column will be read by $T_E X$ shade by '\firstcolumnDSSP' and \secondcolumnDSSP'. The second column is still default.

The HMMTOP algorithm can present its results as plain text or as HTML—plain text needs to be selected here. Further, the output can be formatted in a single line or in an extended form (see the HMMTOP documentation). Both can be read and interpreted by T_EX shade. Importantly, HMMTOP files can contain topology predictions of multiple

sequences. T_EX shade tries to find the correct data based on the respective sequence name. If the sequence name is not found in the file, the first topology data is used. Using an optional parameter (number of the prediction in the file or name) one can define which data from the file is to be used:

$\label{eq:linear} \label{eq:linear} \label{eq:$

PHD predictions: when starting the PHD software do not restrict the prediction to secondary structure or topology alone. This leads to changes in the PHD output file which are not correctly interpretable by T_EX shade due to ambiguities. There is no way around it—thus, run the full prediction.

Now, some information on how T_FX**shade** extracts and displays secondary structure features. In short, it is a two step process. First, T_FXshade analyzes the secondary structure file and extracts all necessary data. This data is converted into a format which is readable and processable by T_{FX} shade using the feature command (see 4.6.3). This command allows one to label sequence stretches graphically. For a detailed explanation see the indicated reference. A list of feature commands is saved in a file with the ending '.sec' for DSSP, STRIDE and PHDsec or '.top' for PHDtopo. Then, in a second step, this file is loaded again and executed. When T_FX<mark>shade</mark> encouters this file a second time, i.e. in a second T_FX run, it uses the already existing file for the output. The great advantage of this method is its flexibility. Due to the simple reason that the feature file can be edited in the meantime. Thus, the user has the ability to change the computer generated file according to his personal needs. On the other hand, one can force T_FX**shade** to write a new file every time by the optional argument [make new] in the include command, e.g. \includePHDsec[make new]{1}{AQP.phd}.

Finally, the appearance of the feature labels can be assigned by the command

Here, $\langle filetype \rangle$ stands for one of the following secondary structure file types: DSSP, STRIDE, PHDsec, PHDtopo or HMMTOP and $\langle type \rangle$ designates the secondary structure type as shown in the right column of the table above. The other arguments $\langle position \rangle$, $\langle labelstyle \rangle$ and $\langle text \rangle$ are almost as described in 4.6.3. One further possibility is to include internal counters for each secondary structure type. Just add one of the following commands to the text in the feature description.

counter	display
\numcount	$1, 2, 3 \dots$
\alphacount	a, b, c
Alphacount	А, В, С
romancount	i, ii, iii
\mathbb{R} omancount	I, II, III

Examples:

\appearance{DSSP}{alpha}{ttop}
 {--->}{\$\alpha\$-helix~\Alphacount}
\appearance{PHDtopo}{TM}{bottom}
 {box[Blue]:TM\numcount[Yellow]}{}

4.7 Displaying and building legends

For each predefined shading mode T_EX shade can print an appropriate legend to explain the used shading colors. The commands \showlegend and \hidelegend display or clear the legend at the end of the alignment. The legend is displayed by default beneath the first residue of the last alignment line. The location can be changed by $\movelegend{\langle x-offset \rangle}{\langle y-offset \rangle}$. Both parameters require a T_EX length, e.g. $\movelegend{5cm}{-2cm}$ moves the legend 5 cm to the right and 2 cm up.

The language for the descriptions is english by default; if the \german.sty package is active legend texts are in german. So far, german, spanish and english are implemented. With the commands \germanlanguage, \spanishlanguage and \englishlanguage switching between the languages is made possible. For the addition of other languages contact me. Finally, the color of the describing legend texts can be set with the command \legendcolor{ $\langle color \rangle$ }.

User defined legends are easily built with the following command $\shadebox{\langle color \rangle}$. Use this command outside the T_EXshade environment, e.g. in the text or in the caption. As $\langle color \rangle$ any color can be designated (see section 5) or one of the following parameters:

- nomatch = the color used for nonmatching residues
- similar = the color used for similar residues
- conserved = the color used for conserved residues

• allmatch = the color used for highly conserved residues (if \allmatchspecial is active)

The command simply prints a shaded box in the specified color then a describing text can be appended. Examples:

```
\shadebox{nomatch}---nonmatching residues
\shadebox{similar}: similar residues
\shadebox{conserved}~conserved residues
\shadebox{Yellow}\quad PKA phosphorylation sites
```

4.8 Adding captions to the alignment

Since T_EX shade v1.5 captions can be added to the alignment. So far, captions were difficult to use when the alignment was bigger than one page and therefore did not fit into a figure environment. The T_EX shade captions behave exactly as normal figure captions. They adopt their style, use the figure counter number and appear in the list of figures as any other figure.

The usage is slightly different from normal captions but intuitive:

The optional $\langle position \rangle$ tells TEX shade to put the caption on top or at the bottom of the alignment. If nothing is stated here the caption will appear at the bottom. The parameter $\langle text \rangle$ just holds the caption text as in the normal \caption. The command can be used at any position within the texshade environment. A simple example would be:

```
\showcaption{A beautiful \TeXshade{} alignment.}
```

4.9 Font handling

4.9.1 Changing font styles

The font styles for the numbering, the sequence names, the sequence residues, the descriptive feature texts and the legends can be changed by several commands.

```
\setfamily{\langle text \rangle}{\langle family \rangle} \setseries{\langle text \rangle}{\langle series \rangle}
```

 $\setshape{\langle text \rangle}{\langle shape \rangle} \setsize{\langle text \rangle}{\langle size \rangle}$

The first parameter selects the text whose style is to be changed. Possible first parameters are numbering, names, residues, features, featurestyles, hideblock, and legend.

The style is set by the second parameter:

command	$\langle 2. parameter \rangle$	
\setfamily	rm	modern roman font family
	sf	sans serif font family
	tt	typewriter font family
\setseries	bf	bold face series
	md	normal series
\setshape	it	italics shape
	sl	slanted shape
	SC	small capitals shape
	up	upright shape
\setsize	tiny	the known T _E X sizes
	scriptsize	
	footnotesize	
	small	
	normalsize	
	large	
	Large	
	LARGE	
	huge	
	Huge	

Example: \setfamily{features}{it} \setseries{features}{bf}

With the command

```
\operatorname{setfont}(\operatorname{text})_{(\operatorname{series})}_{(\operatorname{series})}_{(\operatorname{series})}
```

all four font attributes of one $\langle text \rangle$ can be changed simultaneously. The order of the parameters is as indicated.

Example: \setfont{features}{rm}{it}{bf}{normalsize}

Further, short commands are provided to change single font attributes quickly. The following commands set attributes of feature texts.

```
\featuresrm \featurestiny
\featuressf \featuresscriptsize
```

```
\featurestt \featuresfootnotesize
\featuresbf \featuressmall
\featuresmd \featuresnormalsize
\featuresit \featureslarge
\featuressl \featuresLarge
\featuressc \featuresLARGE
\featuresup \featureshuge
\featuresHuge
```

Corresponding sets are provided for the numbering (\numberingrm etc.), featurestyles (featurestylesrm etc.), names (\namesrm etc.), featurenames (\featurenamesrm etc.), featurestylenames (\featurestylenames etc.), residues (\residuesrm etc.), hideblock labels (hideblockrm etc.), and legend texts (legendrm etc.).

4.9.2 Using PostScript fonts

As already mentioned T_EX **shade** makes intensive use of POSTSCRIPT for shading. Now, that POSTSCRIPT output is active anyway, including POSTSCRIPT fonts is very easy. Just declare in the document header

 $\ensuremath{\scale}\ensuremath$

The typewriter font of TEX is always a topic of discussions. By including the package <code>\usepackage{courier}</code> TEX's typewriter font is replaced by the widely accepted COURIER. Have a look into the directory ..texinputs:latex:psnfss; there, some styles are located which exchange the common TEX fonts by POSTSCRIPT fonts, e.g. avant.sty, bookman.sty, chancery.sty, courier.sty, helvet.sty or utopia.sty. Depending on the style used the <code>\rmdefault-, \sfdefault-, and \ttdefault</code> fonts are substituted partly or completely. Thus, courier.sty for instance exchanges only the type-writer font, whereas bookman.sty sets BOOKMAN as <code>\rmdefault, AVANTGARDE</code> as <code>\sfdefault</code> and COURIER as <code>\ttdefault</code>.

For further information see TOMAS ROKICKI's dvips manual [13].

4.10 Goodies

The following commands give information on sequence properties, such as molecular weight, charge or similarity data. They can be used outside the **texshade** environment directly in the document or in a caption text.

4.10.1 Molweight and charge

During the process of sequence setting T_EX shade sums up the molecular weight and charge of the aligned proteins. This data can be accessed by the following commands.

 $\label{eq:linear} $$ \eqref } \{ \langle Da/kDa \rangle \} \\ charge \{ \langle seqref \rangle \} \{ \langle i/o/N/C \rangle \} \\$

The first parameter $\langle seqref \rangle$ selects the sequence. The second parameter in the \molweight command allows one to switch the units between Dalton (Da) and kilo-Dalton (kDa). The \charge command needs the second parameter for the correct consideration of the charged protein termini. Thus, 'i' refers to internal sequences, 'o' to the overall charge, 'N' to N-terminal sequence parts, and 'C' to the C-terminal end of a protein.

Example: Charge: \charge{1}{o}; Weight: \molweight{1}{Da}

4.10.2 Similarity/identity data and tables

The degree of similarity and identity in percent for any two sequences in the displayed alignment section can be read out with the commands $\percentsimilarity{\langle seqref1 \rangle}{\langle seqref2 \rangle}$ and $\percentidentity{\langle seqref1 \rangle}{\langle seqref2 \rangle}$.

Using the example alignment on page 28 and typing outside the texshade environment in the document text the following phrase:

AQP1 and AQP2 share a sequence similarity of \percentsimilarity{AQP1.pro}{AQP2.pro}\%

will result in the text:

AQP1 and AQP2 share a sequence similarity of 69.6%

Likewise, \percentidentity{1}{2} will give the value 48.4; note that sequences can be referred to by their number or their assigned name. The percent value is calculated by dividing the number of identical or similar positions, respectively, by the total of non-gap positions shared by both sequences. Here, only the part of the alignment is taken into account that is actually displayed. Two residues are considered similar when this is defined by the command \pepsims (see page 46). A full similarity/identity table showing values for all sequences of the alignment can be set using \similaritytable. The labels and number format will be adjusted according to the language settings (4.7).

```
Example (see section 1.6):
   \begin{center}
    \similaritytable
   \end{center}
```

The command generates a valid LATEX tabular environment, which can be embedded into a table environment, e.g.

```
\begin{table}[htdp]
\caption{Text ...}
\begin{center}
   \similaritytable
\end{center}
\end{table}
```

5 The PostScript color selection scheme

POSTSCRIPT provides 64 standard colors. All these colors are predefined in the color.sty. Each color has a pictorial name such as Bittersweet and a distinct composition, e.g. 0% cyan + 75% magenta + 100% yellow + 24% black—the so-called CMYK scheme. TEXshade enhances this color scheme by gray scales in 5% steps. The following colors and grays can be used in TEXshade by simply declaring the name of the color in the respective command, e.g. \consensuscolors:

name	CMYK	name	CMYK
GreenYellow	0.15,0,0.69,0	Yellow	0,0,1,0
Goldenrod	0,0.10,0.84,0	Dandelion	0,0.29,0.84,0
Apricot	0,0.32,0.52,0	Peach	0,0.50,0.70,0
Melon	0,0.46,0.50,0	YellowOrange	0,0.42,1,0
Orange	0,0.61,0.87,0	BurntOrange	0,0.51,1,0
Bittersweet	0,0.75,1,0.24	RedOrange	0,0.77,0.87,0
Mahagony	0,0.85,0.87,0.35	Maroon	0,0.87,0.68,0.32
BrickRed	0,0.89,0.94,0.28	Red	0,1,1,0
OrangeRed	0,1,0.50,0	RubineRed	0,1,0.13,0
WildStrawberry	0,0.96,0.39,0	Salmon	0,0.53,0.38,0
CarnationPink	0,0.63,0,0	Magenta	0,1,0,0
VioletRed	0,0.81,0,0	Rhodamine	0,0.82,0,0
Mulberry	0.34,0.90,0,0.02	RedViolet	0.07,0.90,0,0.34
Fuchsia	0.47,0.91,0,0.08	Lavender	0,0.48,0,0
Thistle	0.12,0.59,0,0	Orchid	0.32,0.64,0,0
DarkOrchid	0.40,0.80,0.20,0	Purple	0.45,0.86,0,0
Plum	0.50,1,0,0	Violet	0.79,0.88,0,0
RoyalPurple	0.75, 0.90, 0, 0	BlueViolet	0.86,0.91,0,0.04
Periwinkle	0.57, 0.55, 0, 0	CadetBlue	0.62, 0.57, 0.23, 0
CornflowerBlue	0.65, 0.13, 0, 0	MidnightBlue	0.98,0.13,0,0.43
NavyBlue	0.94, 0.54, 0, 0	RoyalBlue	1,0.50,0,0
Blue	1,1,0,0	Cerulean	0.94, 0.11, 0, 0
Cyan	1,0,0,0	ProcessBlue	0.96,0,0,0
SkyBlue	0.62, 0, 0.12, 0	Turquoise	0.85, 0, 0.20, 0
TealBlue	0.86, 0, 0.34, 0.02	Aquamarine	0.82, 0, 0.30, 0
BlueGreen	0.85, 0, 0.33, 0	Emerald	1,0,0.50,0
JungleGreen	0.99, 0, 0.52, 0	SeaGreen	0.69, 0, 0.50, 0
Green	1,0,1,0	ForestGreen	0.91,0,0.88,0.12
PineGreen	0.92, 0, 0.59, 0.25	LimeGreen	0.50, 0, 1, 0
YellowGreen	0.44, 0, 0.74, 0	SpringGreen	0.26, 0, 0.76, 0
OliveGreen	0.64, 0, 0.95, 0.40	RawSienna	0, 0.72, 1, 0.45
Sepia	0, 0.83, 1, 0.70	Brown	0, 0.81, 1, 0.60
Tan	0.14, 0.42, 0.56, 0		
White (Gray0)	0,0,0,0	Black (Gray100)	0,0,0,1
Gray5	$0,\!0,\!0,\!0.05$	Gray10	0,0,0,0.10
Gray15	$0,\!0,\!0,\!0.15$	Gray20	0,0,0,0.20

Gray25	0,0,0,0.25	Gray30	0,0,0,0.30
LightGray	0,0,0,0.33	Gray35	$0,\!0,\!0,\!0.35$
Gray40	0,0,0,0.40	Gray45	0,0,0,0.45
Gray50	0,0,0,0.50	Gray	$0,\!0,\!0,\!0.50$
Gray55	0,0,0,0.55	Gray60	$0,\!0,\!0,\!0.60$
Gray65	0,0,0,0.65	DarkGray	$0,\!0,\!0,\!0.66$
Gray70	0,0,0,0.70	Gray75	0,0,0,0.75
Gray80	0,0,0,0.80	Gray85	0,0,0,0.85
Gray90	0,0,0,0.90	Gray95	$0,\!0,\!0,\!0.95$
LightGreenYellow	0.08, 0, 0.35, 0	LightYellow	0,0,0.50,0
LightGoldenrod	0, 0.05, 0.42, 0	LightDandelion	$0,\!0.15,\!0.42,\!0$
LightApricot	0, 0.16, 0.26, 0	LightPeach	$0,\!0.25,\!0.35,\!0$
LightMelon	0, 0.23, 0.25, 0	LightYellowOrange	$0,\!0.21,\!0.50,\!0$
LightOrange	0, 0.31, 0.44, 0	LightBurntOrange	$0,\!0.26,\!0.50,\!0$
LightBittersweet	$0,\!0.38,\!0.50,\!0.12$	LightRedOrange	0, 0.39, 0.44, 0
LightMahagony	0, 0.43, 0.44, 0.18	LightMaroon	0, 0.44, 0.34, 0.16
LightBrickRed	0, 0.45, 0.47, 0.14	LightRed	$0,\!0.50,\!0.50,\!0$
LightOrangeRed	$0,\!0.50,\!0.25,\!0$	LightRubineRed	$0,\!0.50,\!0.07,\!0$
LightWildStrawberry	0,0.48,0.20,0	LightSalmon	0, 0.27, 0.19, 0
LightCarnationPink	0,0.32,0,0	LightMagenta	$0,\!0.50,\!0,\!0$
LightVioletRed	0,0.40,0,0	LightRhodamine	0, 0.41, 0, 0
LightMulberry	0.17, 0.45, 0, 0.01	LightRedViolet	0.04, 0.45, 0, 0.17
LightFuchsia	0.24,0.46,0,0.04	LightLavender	0,0.24,0,0
LightThistle	0.06, 0.30, 0, 0	LightOrchid	0.16, 0.32, 0, 0
LightDarkOrchid	0.20,0.40,0.10,0	LightPurple	0.23, 0.43, 0, 0
LightPlum	0.25, 0.50, 0, 0	LightViolet	0.40, 0.44, 0, 0
LightRoyalPurple	0.38, 0.45, 0, 0	LightBlueViolet	0.43, 0.46, 0, 0.02
LightPeriwinkle	0.29,0.28,0,0	LightCadetBlue	0.31,0.29,0.12,0
LightCornflowerBlue	0.33, 0.07, 0, 0	LightMidnightBlue	0.49, 0.07, 0, 0.22
LightNavyBlue	0.47,0.27,0,0	LightRoyalBlue	0.50, 0.25, 0, 0
LightBlue	0.50, 0.50, 0.0, 0	LightCerulean	0.47, 0.06, 0, 0
LightCyan	0.50, 0, 0, 0	LightProcessBlue	0.48,0,0,0
LightSkyBlue	0.31, 0, 0.06, 0	LightTurquoise	0.43, 0, 0.10, 0
LightTealBlue	0.43,0,0.17,0.01	LightAquamarine	0.41, 0, 0.15, 0
LightBlueGreen	0.43, 0, 0.17, 0	LightEmerald	0.50, 0, 0.25, 0
LightJungleGreen	0.50,0,0.26,0	LightSeaGreen	0.35, 0, 0.25, 0
LightGreen	0.50, 0, 0.50, 0	LightForestGreen	0.46,0,0.44,0.06
LightPineGreen	0.46, 0, 0.30, 0.13	LightLimeGreen	0.25, 0, 0.50, 0
LightYellowGreen	0.22, 0, 0.37, 0	LightSpringGreen	0.13, 0, 0.38, 0
LightOliveGreen	0.32,0,0.48,0.20	LightRawSienna	0, 0.36, 0.50, 0.23
LightSepia	0,0.44,0.50,0.35	LightBrown	0, 0.41, 0.50, 0.30
LightTan	0.07,0.21,0.28,0	0	, , , ,
0		s were derived by divi	ding all values

LightLight- and LightLightLight-versions were derived by dividing all values from Light-color definitions by 2 and 4, respectively.

name	RGB	name	RGB
BlueRed5	0.15, 0.17, 0.55	BlueRed10	0.20, 0.23, 0.57
BlueRed15	0.24, 0.29, 0.60	BlueRed20	0.33, 0.35, 0.64
BlueRed25	0.43, 0.43, 0.68	BlueRed30	0.52, 0.52, 0.73

BlueRed35	0.60,0.60,0.78	BlueRed40	0.70,0.70,0.84
BlueRed45	0.80,0.80,0.85	BlueRed50	0.86,0.82,0.82
BlueRed55	0.87, 0.73, 0.73	BlueRed60	0.89,0.64,0.64
BlueRed65	0.90, 0.55, 0.55	BlueRed70	0.91,0.47,0.46
BlueRed75	0.91, 0.39, 0.37	BlueRed80	0.90,0.33,0.28
BlueRed85	0.89,0.25,0.20	BlueRed90	0.88,0.23,0.14
BlueRed95	0.87,0.21,0.09	BlueRed100	0.87,0.16,0.04
GreenRed5	0,1,0	GreenRed10	0.05, 0.95, 0
GreenRed15	0.10,0.90,0	GreenRed20	0.15, 0.85, 0
GreenRed25	0.20,0.80,0	GreenRed30	0.25, 0.75, 0
GreenRed35	0.30, 0.70, 0	GreenRed40	0.35, 0.65, 0
GreenRed45	0.40, 0.60, 0	GreenRed50	0.45, 0.55, 0
GreenRed55	0.50, 0.50, 0	GreenRed60	0.55, 0.45, 0
GreenRed65	0.60, 0.40, 0	GreenRed70	0.65, 0.35, 0
GreenRed75	0.70, 0.30, 0	GreenRed80	0.75, 0.25, 0
GreenRed85	0.80,0.20,0	GreenRed90	0.85, 0.15, 0
GreenRed95	0.90, 0.10, 0	GreenRed100	0.95, 0.05, 0
ColdHot5	0,0.08,1	ColdHot10	0, 0.29, 1
ColdHot15	0, 0.49, 1	ColdHot20	0, 0.70, 1
ColdHot25	0,0.90,1	ColdHot30	0,1,0.87
ColdHot35	0,1,0.68	ColdHot40	0,1,0.46
ColdHot45	0,1,0.25	ColdHot50	0,1,0.04
ColdHot55	0.16, 1, 0	ColdHot60	0.35, 1, 0
ColdHot65	0.56, 1, 0	ColdHot70	0.79, 1, 0
ColdHot75	0.98, 1, 0	ColdHot80	1,0.82,0
ColdHot85	1,0.60,0	ColdHot90	1,0.40,0
ColdHot95	1,0.20,0	ColdHot100	$0.91,\!0,\!0$
and reverse definition	a DodPluo DodCmo	an HatCald	

and reverse definitions: RedBlue, RedGreen, HotCold.

Type the color names with the upper case letters exactly as described above. For the definition of new colors use one of the color.sty commands:

```
\det\{(name)\} \{ cmyk\} \{ \langle C, M, Y, K \rangle \}
```

```
\det\{ame\} \\ \{g, G, B\} \\
```

The $\langle name \rangle$ can be chosen freely, the values for the color composition must be in the range 0–1, i.e. 0–100% of the respective component ('C' – cyan, 'M' – magenta, 'Y' – yellow, 'K' – black; or 'R' – red, 'G' – green, 'Blue' – blue) separated by commas.

Examples:

```
\label{eq:logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logithtic_logi
```

```
\label{eq:coldHot15}{rgb}{0,0.49,1}
```

6 Listing of the texshade default settings

6.1 Standard definitions

The file texshade.def mirrors all commands which are carried out at the beginning of the texshade environment. Short comments are also included, thus, it is referred to this file for further information.

6.2 Colors used in the different shading modes

Color scheme *blues*:

res.color	shad.color	residues
Black	White	no match
Black	Magenta	similar
White	RoyalBlue	identical
Goldenrod	RoyalPurple	all match

Color scheme greens:

res.color	shad.color	residues
Black	White	no match
Black	GreenYellow	similar
White	PineGreen	identical
YellowOrange	OliveGreen	all match

Color scheme *reds*:

res.color	shad.color	residues
Black	White	no match
Black	YellowOrange	similar
White	BrickRed	identical
YellowGreen	Mahagony	all match

Color scheme grays:

res.color	shad.color	residues
Black	White	no match
Black	LightGray	similar
White	DarkGray	identical
White	Black	all match

Color scheme *black*:

res.color	shad.color	residues
Black	White	no match
Black	White	similar
White	Black	identical
White	Black	all match

Functional mode *charge*:

res.color	shad.color	residues
Black	White	no match
White	Red	acidic
White	Blue	basic

Functional mode *hydropathy*:

res.color	shad. color	residues
Black	White	no match
White	Red	acidic
White	Blue	basic
Black	Yellow	polar uncharged
White	Green	$hydrophobic\ nonpolar$

Functional mode *chemical*:

res.color	shad. color	residues
Black	White	no match
White	Red	acidic
White	Black	aliphatic
White	Gray	aliphatic (small)
White	Green	amide
White	Brown	aromatic
White	Blue	basic
Black	Magenta	hydroxyl
Black	Orange	imino
Black	Yellow	sulfur

Functional mode *rasmol*:

res.color	shad.color	residues
Black	White	no match
Red	White	Asp, Glu
Blue	White	Arg, Lys, His
MidnightBlue	White	Phe, Tyr, Trp
Gray	White	Ala, Gly
Yellow	White	Cys, Met
Orange	White	Ser, Thr
Cyan	White	Asn, Gln
Gree	White	Leu, Val, Ile
Apricot	White	Pro

Functional mode *structure*:

res.color	shad.color	residues
Black	White	no match
Black	Orange	external
Black	Yellow	$\operatorname{ambivalent}$
White	Green	internal

Functional mode *standard area*:

res.color	shad.color	residues
Black	White	no match
Black	BrickRed	G
Black	Orange	A, S
Black	Yellow	C, P
Black	YellowGreen	T, D, V, N
White	PineGreen	I, E
Black	SkyBlue	L, Q, H, M
White	RoyalPurple	F, K
White	RedViolet	Υ
White	Black	R, W

Functional mode *accessible area*:

res.color	shad. color	residues
Black	White	no match
Black	BrickRed	\mathbf{C}
Black	Orange	I, V, G
Black	Yellow	F, L, M, A
Black	YellowGreen	W, S, T, H
White	PineGreen	Р
Black	SkyBlue	Y, D, N
White	RoyalPurple	E, Q
White	RedViolet	R
White	Black	Κ

6.3 Residue weight tables

identity

С	10																			
S	0	10																		
Т	0	0	10																	
Р	0	0	0	10																
А	0	0	0	0	10															
G	0	0	0	0	0	10														
Ν	0	0	0	0	0	0	10													
D	0	0	0	0	0	0	0	10												
Е	0	0	0	0	0	0	0	0	10											
Q	0	0	0	0	0	0	0	0	0	10										
Н	0	0	0	0	0	0	0	0	0	0	10									
R	0	0	0	0	0	0	0	0	0	0	0	10								
Κ	0	0	0	0	0	0	0	0	0	0	0	0	10							
М	0	0	0	0	0	0	0	0	0	0	0	0	0	10						
I	0	0	0	0	0	0	0	0	0	0	0	0	0	0	10					
L	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	10				
V	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	10			
F	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	10		
Y	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	10	
W	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	10
	С	S	Т	Р	A	G	N	D	E	Q	Н	R	K	М	I	L	V	F	Y	W

structural

C S T P A G N D E Q H R K M	10 6 3 5 3 1 0 1 3 0 3 0 3	10 8 8 8 8 5 5 5 5 5 5 5 5	10 6 8 6 5 5 5 5 5 5 5 5 5 5 5 5	10 8 5 5 5 5 3 3 3	10 8 5 6 5 3 3 5 5	10 5 6 3 1 5 3 1	10 8 5 6 5 6 1	10 8 5 3 5 3	10 6 3 5 6 3	10 6 5 6 3	10 6 5 3	10 8 3	10 3	10						
K	0	5	6	3	5	3	6	5	6	6	5	8								
М	3	5	5	3	5	1	1	3	3	3	3	3	3	10						
I	3	3	5	3	3	3	3	1	1	1	3	3	3	6	10					
L	3	3	3	5	3	3	1	1	1	3	5	3	3	8	8	10				
V	3	6	6	6	8	6	3	5	6	3	1	3	5	6	3	3	10			
F	5	5	3	5	5	3	3	1	3	1	3	1	1	5	6	6	6	10		
Y	5	5	3	3	3	3	5	3	1	3	5	1	1	3	5	5	5	8	10	
W	5	3	1	3	3	5	0	0	1	1	1	3	1	5	5	6	5	5	5	10
	С	S	Т	Ρ	A	G	N	D	Е	Q	Н	R	K	М	I	L	v	F	Y	W

PAM250

С	4																			
S	0	3																		
T	-2	1	3																	
P	-3	1	0	6																
А	-2	1	1	1	2															
G	-3	1	0	-1	1	5														
Ν	-4	1	0	-1	0	0	2													
D	-5	0	0	-1	0	1	2	4												
Е	-5	0	0	-1	0	0	1	3	4											
Q	-5	-1	-1	0	0	-1	1	2	2	4										
Н	-3	-1	-1	0	-1	-2	2	1	1	3	6									
R	-4	0	-1	0	-2	-3	0	-1	-1	1	2	6								
Κ	-5	0	0	-1	-1	-2	1	0	0	1	0	3	5							
М	-5	-2	-1	-2	-1	-3	-2	-3	-2	-1	-2	0	0	6						
I	-2	-1	0	-2	-1	-3	-2	-2	-2	-2	-2	-2	-2	2	5					
L	-6	-3	-2	-3	-2	-4	-3	-4	-3	-2	-2	-3	-3	4	2	6				
V	-2	-1	0	-1	0	-1	-2	-2	-2	-2	-2	-2	-2	2	4	2	4			
F	-4	-3	-2	-5	-4	-5	-4	-6	-5	-5	-2	-4	-5	0	1	2	-1	9		
Y	0	-3	-3	-5	-3	-5	-2	-4	-4	-4	0	-4	-4	-2	-1	-1	-2	7	10	
W	-8	-2	-5	-6	-6	-7	-4	-7	-7	-5	-3	2	-3	-4	-5	-2	-6	0	0	17
	С	S	Т	Р	A	G	N	D	E	Q	Н	R	K	М	I	L	V	F	Y	W

PAM100

С	14																			
S	-1	6																		
	_	-	7																	
Т	-5	2	7	10																
Р	-6	1	-1	10	-															
Α	-5	2	2	1	6															
G	-8	1	-3	-3	1	8														
Ν	-8	2	0	-3	-1	-1	7													
D	-11	-1	-2	-4	-1	-1	4	8												
Е	-11	-2	-3	-3	0	-2	1	5	8											
Q	-11	-3	-3	-1	-2	-5	-1	1	4	9										
Н	-6	-4	-5	-2	-5	-7	2	-1	-2	4	11									
R	-6	-1	-4	-2	-5	-8	-3	-6	-5	1	1	10								
Κ	-11	-2	-1	-4	-4	-5	1	-2	-2	-1	-3	3	8							
М	-11	-4	-2	-6	-3	-8	-5	-8	-6	-2	-7	-2	1	13						
I	-5	-4	-1	-6	-3	-7	-4	-6	-5	-5	-7	-4	-4	2	9					
L	-12	-7	-5	-5	-5	-8	-6	-9	-7	-3	-5	-7	-6	4	2	9				
V	-4	-4	-1	-4	0	-4	-5	-6	-5	-5	-6	-6	-6	1	5	1	8			
F	-10	-5	-6	-9	-7	-8	-6	-11	-11	-10	-4	-7	-11	-2	0	0	-5	12		
Y	-2	-6	-6	-11	-6	-11	-3	-9	-7	-9	-1	-10	-10	-8	-4	-5	-6	6	13	
W	-13	-4	-10	-11	-11	-13	-8	-13	-14	-11	-7	1	-9	-11	-12	-7	-14	-2	-2	19
	10	-	10			10	Ŭ	10			•	-	U			•		-	-	-0
	С	S	Т	Р	А	G	Ν	D	E	Q	Н	R	K	М	I	L	V	F	Y	W

BLOSUM62

С	9																			
S	-1	4																		
T	-1	1	4																	
P	-3	-1	4 1	7																
-			-	•																
A	0	1	-1	-1	4															
G	-3	0	1	-2	0	6														
Ν	-3	1	0	-2	-2	0	6													
D	-3	0	1	-1	-2	-1	1	6												
Е	-4	0	0	-1	-1	-2	0	2	5											
Q	-3	0	0	-1	-1	-2	0	0	2	5										
Н	-3	-1	0	-2	-2	-2	1	1	0	0	8									
R	-3	-1	-1	-2	-1	-2	0	-2	0	1	0	5								
Κ	-3	0	0	-1	-1	-2	0	-1	1	1	-1	2	5							
М	-1	-1	-1	-2	-1	-3	-2	-3	-2	0	-2	-1	-1	5						
I	-1	-2	-2	-3	-1	-4	-3	-3	-3	-3	-3	-3	-3	1	4					
L	-1	-2	-2	-3	-1	-4	-3	-4	-3	-2	-3	-2	-2	2	2	4				
V	-1	-2	-2	-2	0	-3	-3	-3	-2	-2	-3	-3	-2	1	3	1	4			
F	-2	-2	-2	-4	-2	-3	-3	-3	-3	-3	-1	-3	-3	0	0	0	-1	6		
Y	-2	-2	-2	-3	-2	-3	-2	-3	-2	-1	2	-2	-2	-1	-1	-1	-1	3	7	
W	-2	-3	-3	-4	-3	-2	-4	-4	-3	-2	-2	-3	-3	-1	-3	-2	-3	1	2	11
	С	S	Т	Р	А	G	Ν	D	E	Q	Н	R	K	М	Ι	L	V	F	Y	W

7 Quick Reference

The T_EXshade logo

\TeXshade

The T_EX_{shade} environment (15 ff.)

 $\label{eq:lasshade} [\langle parameterfile \rangle] \{ \langle alignmentfile \rangle \} \\ further T_EX shade commands, if needed \\ \end{texshade} \}$

Predefined shading modes

$seqtype{\langle type \rangle}$	(P - peptide, N - nucleotide)	[41]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	${ mode } $	[44]

$\langle mode \rangle$	$\langle option \rangle$	
identical	allmatchspecial/number	
similar	allmatchspecial/number	
T-Coffee	$\langle filename \rangle$	
diverse	$\langle seqref \rangle$	
functional	$\langle type \rangle$	charge
		hydropathy
		structure
		chemical
		rasmol
		standard area
		accessible area

$\ \[\langle percentage \rangle]$	[44]
\shadeallresidues	[49]

Shading colors $(44 \, \text{ff.})$

```
\label{eq:shadingcolors} $$ (blues, reds, greens, grays, black) \\ nomatchresidues{$ (res. col.)}{$ (shad. col.)}{$ (case)}{$ (style)} \\ similarresidues{$ (res. col.)}{$ (shad. col.)}{$ (case)}{$ (style)} \\ conserved residues{$ (res. col.)}{$ (shad. col.)}{$ (case)}{$ (style)} \\ allmatchresidues{$ (res. col.)}{$ (shad. col.)}{$ (case)}{$ (style)} \\ defshadingcolors{$ (name)}} $
```

 $\label{eq:linearized_linearized$

Residue grouping

$pepsims{\langle residue \rangle}{\langle similars \rangle}$	[46]
$pepgroups{\langle group1 \rangle, \langle group2 \rangle, \ldots, \langle groupn \rangle}$	[46]
$DNAsims{\langle residue \rangle}{\langle similars \rangle}$	[46]
$DNAgroups{\langle group1 \rangle, \langle group2 \rangle, \ldots, \langle groupn \rangle}$	[46]

Definition of new functional shading modes

\clearfuncgroups	[49]
$\timescale \timescale \timescal$	
$ \{ \langle case \rangle \} \{ \langle style \rangle \} $	[50]

Appearance of the consensus line

	[[1]
$\theta = \frac{1}{2} \left\{ \left\langle percentage \right\rangle \right\}$	[51]
$\operatorname{constosingleseq} \{ \langle seqref \rangle \}$	[51]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[51]
$\exp [\langle filename \rangle] \{\langle seqref \rangle\}$	[51]
\hideconsensus	[51]
$nameconsensus{\langle name \rangle}$	[51]
$\det consensus (symbol 1) \{ (symbol 2) \} \{ (symbol 3) \}$	[52]
$\consensus colors{\langle res. col. 1 \rangle}{\langle shad. col. 1 \rangle}$	
$\{\langle res. col. 2 \rangle\}\{\langle shad. col. 2 \rangle\}$	
$\{\langle res. col. 3 \rangle\}$	[52]

$weighttable{ (table) }$	[53]
$setweight{\langle res.1 \rangle}{\langle res.2 \rangle}{\langle value \rangle}$	[54]
$gappenalty{\langle value \rangle}$	[54]

Sequence logos

$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[54]
\hidesequencelogo	[54]
$\clearlogocolors[\langle color \rangle]$	[55]
$\log color{\langle residues \rangle}{\langle color \rangle}$	[55]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[55]

\hidelogoscale	[55]
$logostretch{\langle factor \rangle}$	[55]
$namesequencelogo{\langle name \rangle}$	[55]
\dofrequencycorrection	[55]
\undofrequencycorrection	[55]

Subfamily logos

$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[56]
\hidesubfamilylogo	[56]
$setsubfamily{\langle seqrefs \rangle}$	[56]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[56]
\hidenegatives	[56]
$\mbox{namesubfamilylogo}[\langle neg.name \rangle] \{\langle name \rangle\}$	[56]
$\columnwidth{\columnwidth$	[56]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[56]
\hiderelevance	[56]

Appearance of the sequence lines

$\ \ \ \ \ \ \ \ \ \ \ \ \ $	[56]
$\ \ \ \ \ \ \ \ \ \ \ \ \ $	[56]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[57]
$\namescolor{\langle color \rangle}$	[57]
$\namecolor{\langle seq1 \rangle, \ldots, \langle seq n \rangle}{\langle color \rangle}$	[57]
\hidenames	[57]
$\lambda \{seq1\}, \ldots, (seq n)\}$	[57]
$\operatorname{\mathbb{C}}^{\operatorname{\mathbb{C}}}$	[57]
$\operatorname{numbercolor}(\operatorname{seq}1), \ldots, \operatorname{seq}n) \in \operatorname{color})$	[57]
\hidenumbering	[57]
$ \left(seq1 \right), \ldots, \left(seq n \right) $	[57]
\hideresidues	[61]
\showresidues	[61]
\times	[57]
\allowzero	[57]
\disallowzero	[57]
$\ensuremath{seqref}{eq:eq:eq:eq:eq:eq:eq:eq:eq:eq:eq:eq:eq:e$	[58]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[58]
$\operatorname{rulersteps} \{ \langle number \rangle \}$	[58]
$\operatorname{color}{\langle color \rangle}$	[58]
\hideruler	[58]

\rotateruler	[58]
\unrotateruler	[58]
$\mbox{namerulerpos}{\langle number \rangle}{\langle text \rangle}[\langle color \rangle]}$	[58]
$\gapchar{\langle symbol \rangle}$ (incl. rule)	[58]
$\gapcolors{\langle symbol \ color \rangle}{\langle background \ color \rangle}$	[58]
\showleadinggaps	[59]
\hideleadinggaps	[59]
$\t \in \{symbol\}$	[59]
$fingerprint{ (res. per line)}$	[63]

Displaying selected residues in the alignment

$\setends[\langle startnumber \rangle] \{\langle seqref \rangle\} \{\langle startstop \rangle\}$	[59]
$\operatorname{setdomain} \{\langle seqref \rangle\} \{\langle selection \rangle\} (see \ baderegion p. 64)$	[60]
$\operatorname{domaingaprule}{\langle thickness \rangle}$	[60]
$\boldsymbol{\boldsymbol{\boldsymbol{\boldsymbol{\boldsymbol{\boldsymbol{\nabla}}}}}}$	[60]

Hiding, killing, separating and ordering

$\label{eq:linear} \$	[61]
\hideseqs	[61]
\showseqs	[61]
$ \left(seq1 \right), \left(seq2 \right), \dots, \left(seq \ n \right) $	[61]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[61]
$separationline{\langle seqref \rangle}$	[61]
\smallsep	[61]
\medsep	[61]
\bigsep	[61]
$vsepspace{\langle length \rangle}$	[61]
$\operatorname{Seq1}, \operatorname{Seq2}, \ldots, \operatorname{Seq}n$	[62]

Residues per line and further settings

$\ensuremath{residuesperline}\$	[62]
$\ensuremath{residuesperline}{\operatorname{sumber}}$	[62]
$\operatorname{Charstretch}{\langle factor \rangle}$	[62]
$\linestretch{\langle factor \rangle}$	[62]
$\operatorname{\mathbb{R}}^{\operatorname{\mathbb{R}}}$	[62]
\smallblockskip	[63]
\medblockskip	[63]
\bigblockskip	[63]

\noblockskip	[63]
$vblockspace{\langle length \rangle}$	[63]
\flexblockspace	[63]
\fixblockspace	[63]
$\ \ \ \ \ \ \ \ \ \ \ \ \ $	[63]

Individual shading and labeling of sequence stretches

```
\label{eq:selection} \end{tabular} $$ \eqref \end{tabular} $$ \eqref \end{tabular} $$ \eqref \end{tabular} $$ \eqref \eqre \eqref \eqref \eqref \eqref \eqref \eqre \eqr
```

[65] $\changeshadingcolors{\langle seqref \rangle}{\langle selection \rangle}{\langle name \rangle}$ [66] $\ensuremath{\mathsf{emphregion}}{\langle seqref \rangle} \{\langle selection \rangle\}$ [66] $\mathbb{C} \left(seqref \right)$ [66][66] $\ensuremath{\mathsf{emphdefault}}{\langle style \rangle}$ $\operatorname{tintregion} \{\langle seqref \rangle\} \{\langle selection \rangle\}$ [66] $\tilde{\langle seqref \rangle} \{\langle selection \rangle\}$ [66] $\tilde{\langle effect \rangle}$ weak, normal, strong [66] $\lowerregion{\langle seqref \rangle}{\langle selection \rangle}$ [66] $\lowerblock{\langle seqref \rangle}{\langle selection \rangle}$ [66] $frameblock{\langle seqref \rangle}{\langle selection \rangle}{\langle color \rangle}{\langle length \rangle}}$ [67] $feature{\langle position \rangle} \{\langle seqref \rangle\} \{\langle selection \rangle\}$

```
\{\langle labelstyle \rangle\}\{\langle text \rangle\} [67]
```

 $= \{\texttt{translate} \lfloor \langle color \rangle \rfloor \}$

 $= \{ bar[\langle min \rangle, \langle max \rangle] : \\ \langle properties/file/data \rangle[\langle color(, bgcolor) \rangle] \} \\ = \{ color[\langle min \rangle, \langle max \rangle] : \\ \langle properties/file/data \rangle[\langle scale \rangle] \} \\ \langle properties \rangle: hydrophobicity, charge, \\ molweight, conservation \\ \langle scale \rangle: Gray, BlueRed, RedBlue, \\ GreenRed, RedGreen, ColdHot, \\ HotCold, T-Coffee \\ \end{cases}$

$ttopspace{\langle length \rangle}$	[67]
$\topspace{\langle length \rangle}$	[67]
$bottomspace{\langle length \rangle}$	[67]
$bbottomspace{\langle length \rangle}$	[67]
$featurerule{\langle length \rangle}$	[69]
$bargraphstretch{\langle factor \rangle}$	[72]
$colorscalestretch{\langle factor \rangle}$	[72]
$ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[70]
$geneticcode{\langle filename \rangle}$	[70]
$backtranslabel[\langle size \rangle] \{\langle style \rangle\}$	[71]
$backtranstext[\langle size \rangle] \{\langle style \rangle\}$	[73]

$$\{\langle style \rangle\} = \{$$
horizontal $\}$
= $\{$ alternating $\}$
= $\{$ zigzag $\}$
= $\{$ oblique $\}$
= $\{$ vertical $\}$

$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[73]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[73]
$\tilde{ttttopbbbbottom}$	[73]
$\theta \in \{\langle ttttopbbbbottom \rangle\}$	[73]
\hidefeaturenames	[73]
\hidefeaturestylenames	[73]
$featurenamescolor{color}$	[74]
$featurestylenamescolor{color}$	[74]
$featurenamecolor{\langle ttttopbbbbottom \rangle}{\langle color \rangle}$	[74]
$featurestylenamecolor{\langle ttttopbbbbottom \rangle}{\langle color \rangle}$	[74]

Including secondary structure information

$\label{eq:linear} \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \$	[75]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[75]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[75]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[75]
$\label{eq:linear} \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \$	[75]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[76]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[76]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[76]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[76]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[76]
$\tilde{structures}$	[76]
$\tilde{structures}$	[76]
$\tilde{structures}$	[76]
$\tilde{\phi} $	[76]
$\tilde{structures}$	[76]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[77]
\numcount	[77]
\alphacount	[77]
\Alphacount	[77]
\romancount	[77]
\Romancount	[77]
\firstcolumnDSSP	[76]
\secondcolumnDSSP	[76]

Displaying and building legends

\showlegend	[78]
\hidelegend	[78]
$movelegend{\langle x-offset \rangle}{\langle y-offset \rangle}$	[78]
\germanlanguage, \spanishlanguage, \englishlanguage	[78]
$\ensuremath{\label{color}}$	[78]
$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[78]

Adding captions to the alignment

$\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	[79]
$\ \ \ \ \ \ \ \ \ \ \ \ \ $	[79]

Font handling

```
setfamily{\langle text \rangle}{\langle family \rangle}
                                                                                  [79]
setseries{\langle text \rangle}{\langle series \rangle}
                                                                                  [79]
setshape{\langle text \rangle}{\langle shape \rangle}
                                                                                  [79]
setsize{\langle text \rangle}{\langle size \rangle}
                                                                                  [79]
\operatorname{setfont}(\langle text \rangle) \{\langle family \rangle\} \{\langle series \rangle\} \{\langle shape \rangle\} \{\langle size \rangle\}
                                                                                  [80]
                                                                                  [80]
\featuresrm
                   \featurestiny
\featuressf
                    \featuresscriptsize
\featurestt
                    \featuresfootnotesize
\featuresbf
                    \featuressmall
\featuresmd
                    \featuresnormalsize
\featuresit
                     \featureslarge
\featuress1
                    \featuresLarge
                    \featuresLARGE
\featuressc
\featuresup
                    \featureshuge
                     \featuresHuge
```

Corresponding sets are provided for the numbering (\numberingrm etc.), featurestyles (featurestylesrm etc.), names (\namesrm etc.), featurenames (\featurenamesrm etc.), featurestylenames (\featurestylenames etc.), residues (\residuesrm etc.), hideblock labels (hideblockrm etc.), and legend texts (legendrm etc.).

Goodies—molweight and charge

$\mathbf{e}_{\langle ba/kDa \rangle} $	[82]
$\ensuremath{\label{eq:charge}} \langle eqref \rangle \} \{ \langle i/o/N/C \rangle \}$	[82]
$percentsimilarity{(seqref1)}{(seqref2)}$	[82]
$percentidentity{(seqref1)}{(seqref2)}$	[82]
\similaritytable	[83]

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