

# MoltenProt Documentation

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## Table of Contents

1.	What is MoltenProt?	1
2.	Step-by-step instructions	2
3.	FAQ	3
3.1.	General questions	3
3.2.	Using MoltenProt	3
4.	Graphical user interface	4
4.1.	Main window	4
4.2.	Toolbar	4
4.3.	File Menu	5
4.4.	Actions   Analysis	5
4.5.	Actions   Edit layout	7
4.6.	Actions   Select / Deselect	7
4.7.	Actions   Settings	7
5.	Command-line interface	9
6.	Input/Output	10
6.1.	Supported formats	10
6.2.	Output files	11
7.	Models	13
7.1.	Overview	13
7.2.	Extensive and intensive readouts	13
7.3.	Equilibrium models	13
7.4.	Empirical models	15
7.5.	Kinetic models	15
8.	Credits & copyright	17
8.1.	MoltenProt	17
8.2.	Dependencies	17
9.	References	22

## 1. What is MoltenProt?

MoltenProt is a program to fit sigmoidal curves obtained with label-free protein unfolding assays, such as NanoDSF or circular dichroism measurements [1]. In addition to widely-used melting temperature ( $T_m$ ) MoltenProt uses other curve characteristics to rank the results in terms of their (thermo)stability.

Furthermore, MoltenProt offers a panel of protein unfolding models, including equilibrium unfolding, irreversible unfolding and the Lumry-Eyring model. See [Models](#) for more information.

MoltenProt provides a [GUI](#) for exploratory data analysis and a [CLI](#) for batch-processing.

## 2. Step-by-step instructions

1. Obtain a dataset, where protein unfolding is monitored with a label-free technique as a function of temperature.  
**NOTE:** to load a demo dataset from the GUI use File > Load sample data
2. If the input data is in XLSX format, use a spreadsheet editor to annotate the samples (sheet "Overview").
3. Start the MoltenProt GUI and load the dataset. Hover on individual samples with mouse to view the raw curves. Click on wells to display several curves. If multiple datasets are present in the file, a combobox will appear in the toolbar. Inspect the curves. If needed, set bad curve annotation to "Ignore" using Layout editor.
4. Open the analysis menu and select the model for each dataset. The default settings usually provide the best performance of the fit, however, the model may not reflect the real nature of protein unfolding.
  - 4.1. If curves contain spikes, they can be removed by trimming some values in the beginning or end of the curve. In more difficult cases, a median filter can be applied to smooth out the spikes.
  - 4.2. If curves are too noisy, the signal strength may be improved by averaging datapoints to a larger degree step (shrinking).
5. Once the analysis is done, the sample stability will be color-coded on a heatmap. By default the model-supplied ranking parameter will be used for the heatmap. Other useful parameters for heatmap coloring will be available in a combobox. Click on the samples of interest to compare their fit curves side-by-side and show the fit parameters in a table.
6. Inspect highest/lowest ranked curves. How noisy is the measurement? Does the fit result reflect the curve features?
  - 6.1. If a particularly noisy curve distorts the heatmap, it can be removed from analysis by annotating the sample as "Ignore" in Layout editor and re-running analysis.
  - 6.2. If needed, perform fine-tuning of curve fitting parameters.
  - 6.3. To store all analysis and visualization settings save a MoltenProt session in JSON format. The session file can be loaded later for re-analysis or data exporting.
7. Export the data using the format that is most appropriate for downstream analysis.

## 3. FAQ

### 3.1. General questions

#### 3.1.1. Is MoltenProt suitable for analysis of assays employing fluorescent dyes to report protein unfolding?

No, MoltenProt is written for label-free assays, such as tryptophan fluorescence and circular dichroism. Dye-based unfolding assays exhibit highly non-linear post-transition baselines, which are more difficult to model, and overall quality of the data is usually inferior.

#### 3.1.2. Why is (Nano)DSF an inappropriate term?

DSF stands for "differential scanning fluorimetry" and was coined as an analogy to DSC, differential scanning calorimetry. In DSC one measures *difference* in heat capacity relative to a reference cell, while in unfolding assays with fluorescence readouts there is no such reference. Thus, they should not be called "differential". In MoltenProt documentation this technique is referred to as "label-free protein unfolding assay".

## 3.2. Using MoltenProt

#### 3.2.1. How can I select a subset of samples?

Open the [layout dialog](#) and set unneeded samples to "Ignore".

#### 3.2.2. I cannot open my file, what shall I do?

First, check if correct file [import settings](#) were specified. Second, compare your file with the provided demo data, in particular, for XLSX files the correct naming of sheets is important.

#### 3.2.3. Why is the temperature scale in Kelvins? It is annoying!

*Official answer:* Kelvins are the SI units for temperature.

*Unofficial answer:* to avoid bugs in thermodynamic calculations.

#### 3.2.4. Can I process more than 96 samples?

No, datasets with more than 96 samples have to be split into smaller chunks.

#### 3.2.5. Why derivative curves are not exported?

Computation of a derivative for experimental data is a tricky task and inevitably introduces noise, so it is not recommended for precise estimates of Tm. On the other hand, smoothed derivative curves proved useful in visual inspection of data, so MoltenProt can display them in the plot window.

### 3.2.6. I am not happy with the fit results, what should I try?

If the baselines are not adequately fit, increase baseline\_stdev and/or trim 5-10 degrees in the beginning or end of the curve (see [Preprocessing](#)).

### 3.2.7. My data contains spikes (e.g. from a bubble), is it bad?

Short spikes (5 degrees wide or so) usually pose no problem to the fit, however, such curves will have a higher S (standard error of the estimate). If a spike is very close to the beginning or end of the curve, consider removing it by trimming, because it will interfere with baseline pre-estimation. Spikes can be also removed with [median filtering](#), however, this distorts the experimental signal.

### 3.2.8. What is the advantage of using the "[irrev](#)" model?

This model assumes that protein unfolds irreversibly, which is a more realistic assumption compared to the classic thermodynamic model, where full reversibility of unfolding is assumed. Indeed, how often have you seen proteins that remain functional after being boiled? Also, irrev model enables comparison of data collected at different scan rates. In terms of relative result ranking, however, we did not observe any differences between irrev and [classic thermodynamic models](#) (e.g. santoro1988, see [2]).

## 4. Graphical user interface

### 4.1. Main window



- 1. Heatmap panel:** samples are color-coded with the selected curve characteristics. Hover-on with a mouse to visualize the data in the Plot window. Click several samples to visualize them side-by-side on the Plot window; their fit parameters will be shown in the Result table.

2. **Result table**: displays characteristics of selected curves. The set of characteristics to be displayed depends on the type of analysis performed.
3. **Plot window**: visualizes data requested by the user. A variety of plots can be displayed; use [Settings](#) for fine tuning.
4. **Protocol window**: displays the log of the data analysis including informational messages and warnings. Errors in analysis produce a pop-up window.
5. **Toolbar** and **menus** provide access to the functions of MoltenProt; **window decorator** is managed by the operating system.

## 4.2. Toolbar

### 4.2.1. Readout combobox

Switches between the readouts present in the input file, e.g. F330, F350 and Ratio. If the input file contains a single readout (e.g. plain CSV), then Readout combobox will not be shown.

### 4.2.2. Heatmap combobox

Selects a curve characteristic to color the heatmap in the GUI. Available options depend on the chosen analysis [model](#).

### 4.2.3. Font settings

Loads the menu to adjust font size, type, etc in MoltenProt. Useful for scaling the program window on high-resolution displays.

## 4.3. File Menu

### 4.3.1. File | New

Start a new MoltenProt session.

### 4.3.2. File | Load

Open one of the [supported file formats](#): comma-separated values (CSV), NanoDSF processed data (XLSX) or MoltenProt session (JSON).

### 4.3.3. File | Load sample data

Opens the folder with demo data (in all supported formats).

### 4.3.4. File | Export

Export results with selected settings to a directory.

#### 4.3.5. File | Save as JSON

Save the current MoltenProt session.

#### 4.3.6. File | Quit

Terminate the program.

### 4.4. Actions | Analysis

Set analysis settings and process data. OK button will run the analysis, Cancel button will close the window, Reset to defaults button will supply default values to all analysis parameters.

#### 4.4.1. Basic settings

This tab displays a table with available datasets (1 in case of CSV, up to 5 in case of XLSX input file type) and a combobox with possible models of analysis:

1. **santoro1988**: fast and robust fitting based on equilibrium two-state unfolding model;
2. **santoro1988i**: same as 1, but with an additional unfolding intermediate (three-state model);
3. **santoro1988d**: fast, but less robust fitting, which works descriptively, i.e. not assuming any unfolding mechanism; the idea is to provide the best quantitative description of the experimental curve;
4. **santoro1988di**: same as 3, but suitable for fitting unfolding curves with one intermediate (two peaks observed in the derivative plot);
5. **irrev**: irreversible two-state unfolding; protein unfolding is described kinetically, rather than using thermodynamics. This is a very common case, because most proteins do not reach equilibrium and unfold irreversibly. Computation requires numeric integration, so data processing is slow;
6. **lumry\_eyring**: Lumry-Eyring model for protein unfolding coupled with aggregation; can be used only if Scattering is available in the input data. First, the kinetics of aggregation are estimated using irrev model, and then unfolding and refolding kinetics are estimated. Computation requires numeric integration, so data processing is slow;
7. **skip**: the dataset will not be processed at all. The raw data will be preserved and can be re-analysed later on.

For more details on the models implemented in MoltenProt see [Models](#). In most cases the choice of the model is the only decision required from the user.

#### 4.4.2. Pre-processing

- **Median filtering:** remove spikes from the data by applying a median filter. The window size, i.e. the number of datapoints used to compute the median, is specified in temperature degrees. Median filter removes information from the data, and curve fitting is usually robust and not affected by spikes, so this option is rarely needed.
- **Shrink data:** shrink data to a specified degree step. This step removes information from the data and decreases certainty of the fit, but may help expose global trends in the data. Also, shrunk data are processed faster.
- **Remove from curve start/end:** drop some datapoints in the beginning of the end of the curve. This option may be helpful if the signal spikes at the start or end of the experiment.

#### 4.4.3. Misc. options

- **Savitzky-Golay window size:** window size for Savitzky-Golay filter to calculate the smoothed first derivative. The window size is specified in temperature degrees and converted to an odd number of datapoints automatically. The smoothed derivative is used in data visualization and also provides the initial value for Tm.
- **Data length for baseline estimation:** how many degrees in the beginning and the end of the curve are pure baselines, i.e. temperature dependence of the signal with 0% and 100% protein molecules unfolded. The stretches of the data will be used to generate initial values for baseline fit parameters.
- **Baseline bounds (n\*stdev):** after initial baseline estimation (see previous option), MoltenProt uses the standard deviations for the slope and the intercept as the parameter bounds for pre- and post-transition baselines in the fitting of the full unfolding curve. In problematic cases this prevents the baselines from moving too far away from the experimental data.
- **Heat capacity change ( $\Delta C_p$ ):** provide the value in J/mol/K, which will be used to extrapolate  $\Delta G_u$  from the unfolding region (around Tm) to the standard temperature of 298.15 K. If  $\Delta C_p$  is zero, then the extrapolation will be linear. For soluble proteins of 30 kDa and below  $\Delta C_p$  can be estimated by multiplying the number of residues by 58 [3]. This value is only relevant for models based on equilibrium unfolding.

### 4.5. Actions | Edit layout

Create or edit annotations of individual samples. This information will be shown in the GUI and, where applicable, written to the output files. Annotations can be

added manually via the opened layout dialog or loaded from a CSV file. For XLSX files the recommended way to supply annotations is to edit the "Overview" sheet in the original XLSX file with a spreadsheet editor. See [Supported Formats](#) for more info.

All changes made to the layout can be undone using the Reset button.

Context menu for the layout dialog (right mouse button) provides additional options:

- **Blank:** mark selected samples as blank buffer. During the analysis these curves will be averaged and subtracted from all other curves. This can be used to remove signal of the buffer. Note that the proper blank subtraction for F330/F350 Ratio data is not implemented.
- **Ignore:** skip selected samples in the analysis.
- **Clear selected cells:** clear any annotations in the selected cells.

## 4.6. Actions | Select / Deselect

Display/hide all curves in the dataset.

## 4.7. Actions | Settings

### 4.7.1. Import

- **CSV:** parameters for parsing unfolding data in CSV format:
  - Separator: character that separates data entries.
  - Decimal separator: character that indicates the decimal digit.
  - Denaturation: indicate if the temperature scale in the input file is in Kelvins or Celsius.
  - Scan rate: heating rate in degrees/min. Scan rate is relevant only [for non-equilibrium models](#).
  - Spectrum: indicate if data comes from a spectrum acquisition. The first column must contain the temperature values, and all other columns must correspond to the signal at a specific wavelength. The wavelength must be indicated in the first row.
- **XLSX:** parameters for parsing XLSX data (NanoDSF):
  - Refolding data: Indicate if the refolding ramp was used in the experiment; this is needed for correct parsing of input files. The refolding data is treated independently from the unfolding data. For fully reversible folding both curves must have similar fit parameters.
  - Raw data: Indicate if “raw” data was exported from the NanoDSF machine. In this data each sample has its own time scale.

## 4.7.2. Export

- **Data format:**
  - "None": create no output tables.
  - "CSV": output comma-separated files with UTF encoding and each table will be an individual file.
  - "XLSX": export a single file with multiple sheets.
- **Report format:**
  - "None": no reports are generated.
  - "PDF": a comprehensive PDF report with tabulated data, heatmap and plots of individual samples. A separate file will be created for each readout.
  - "Summary (XLSX)": produce an XLSX file containing a result table for all readouts, but no curves or images will be generated.
  - "Interactive (HTML)": the report will contain a full package of data including XLSX files with raw, fit, baseline-corrected curves, analysis results and standard deviations. The files will be linked in a single HTML file, which also allows viewing plots of individual samples. The main advantage of HTML reports is that they do not require installation of MoltenProt and can be viewed in any modern web-browser.

For more details see [Output files](#).

## 4.7.3. Misc

- **Parallel processes:** MoltenProt will run some steps with the specified number of subprocesses. This can speed up the workflow, but will also consume more computer resources.
- **Colormap for heatmap:** Choose one of the matplotlib colormaps for the main heatmap:



Colormaps with "\_r" will have inverse color direction.

## 4.7.4. Plots

- **Display curve:**
  - "Experimental signal": show raw experimental data.

- "Baseline-corrected": pre- and post-transition baselines are subtracted from the data to get a sigmoidal curve ranging from 0 to 1. This viewing mode simplifies comparison between samples.
- **Display as:**
  - "Datapoints + Fit": show both experimental and the fit result.
  - "Datapoints": only experimental data.
  - "Fit": only fit data.
- **Use heatmap color:** if enabled, than the colors from the buttons in the heatmap panel will be applied to respective curves. Otherwise a random color will be given to every plotted curve.
- **Baselines:** show pre- and post-transition baselines obtained in the fitting as dashed lines. If one of the baselines is not visible or is too far from the data, then the fitting may have gone wrong.
- **Vertical lines:** show characteristic temperatures (e.g. T<sub>m</sub> or T<sub>onset</sub>) as vertical lines on the plot.
- **Show every:** Show only n-th experimental datapoint. This is useful to unclutter dense experimental curves.
- **Derivative plot:** Create an additional plot showing the smoothed first derivative of experimental data.
- **Legend:**
  - "None": do not show legend.
  - "ID": show sample ID in the legend (A1-H12).
  - "Annotation": show the sample annotation (as recorded in the [layout](#)).

## 5. Command-line interface

Upon installation, MoltenProt is accessible as a Python module:

```
python -m moltenprot --help
```

All options implemented in the GUI are also available via the CLI. The only difference is that per-readout model settings are not available. The main usage for CLI is to perform processing of multiple datasets with the same parameters:

```
python -m moltenprot -i dataset1.xlsx dataset2.xlsx -o all_datasets -j 2 --exclude_readout 330nm --model_sct santoro1988d --report pdf
```

This command will process files dataset1.xlsx and dataset2.xlsx, write the output (PDF report) to folder all\_datasets. The readout called "330nm" will be excluded and for scattering data model santoro1988d will be applied. All other

datasets will be processed with default model santoro1988. The program will run in two parallel processes.

## 6. Input/Output

### 6.1. Supported formats

#### 6.1.1. CSV

Comma-separated value (CSV) file for MoltenProt must follow several rules:

- First row contains column names.
- One column is called "Temperature" and contains the X-axis values.
- All other columns have an alphanumeric index similar to a 96-well plate (from A1 to H12).

```
Temperature,A1,A2, ... ,H12
20,1300,1500,...,1600
21,1400,1600,...,1700
...
95,2000,2100,...,2200
```

Under the hood MoltenProt uses pandas.read\_csv to parse CSV files, so any separator supported by this module can be used in MoltenProt.

#### 6.1.2. Layout CSV

These files provide a description of individual samples and can be added to MoltenProt session with [Layout editor](#). Files should follow several requirements:

- First row contains column names.
- One column is called "ID" and contains an alphanumeric index similar to a 96-well plate (from A1 to H12).
- One column is called "Condition" and contains the annotations.
- The file can be only comma-separated, any other separators are not allowed; it is also recommended that all text is quoted.
- An optional column can be called "dCp" and can contain per-sample values of heat capacity change.

```
"ID","Condition"
"A1","Ultrapure water"
"A2","Original buffer"
...
"H12","Blank"
```

### 6.1.3. Spectrum CSV

The file formatting is similar to standard CSV, however, columns instead of alphanumeric IDs can be named after the wavelengths in the recorded spectrum. For instance:

```
Temperature,220nm,221nm, ...,500nm  
20,1300,1500,...,1600  
21,1400,1600,...,1700  
...  
95,2000,2100,...,2200
```

If there are more than 96 columns, then excess columns will be skipped at a regular interval (e.g. every 5<sup>th</sup> will be dropped).

### 6.1.4. XLSX

XLSX files with "processed" or "raw" data generated by PR.ThermControl (NanoTemper Technologies GmbH, tested with v. 2.1.2 and v. 2.3.1) can be opened directly in MoltenProt. The annotations provided in the "Overview" sheet are imported as well. While MoltenProt offers basic capabilities for editing annotations, it is recommended to set all annotations in the "Overview" sheet using a full-featured spreadsheet editor.

**NOTE:** An additional readout, F350-F330 (deltaF), is computed automatically. For more info see [Models](#) section.

### 6.1.5. JSON

MoltenProt uses JavaScript Object Notation (JSON) format to store sessions, i.e. the current state of the program. Sessions contain raw and processed data as well as annotations, analysis options used, timestamps etc, thus providing an easy way to save results for later viewing.

## 6.2. Output files

File export settings can be adjusted in [Settings|Export](#). MoltenProt usually writes out multiple files, so it is recommended to export results to a dedicated directory. The exception is saving of JSON sessions, where everything is written into a single file.

### 6.2.1. Curves

For each readout present in the input file (e.g. F330, F350, Ratio) CSV export will produce a separate file for each curve type (see below). XLSX export will create a single XLSX file for each readout present in the input file. The following curve types are available:

- **Raw data** (not exported to CSV): original data read from the input file.
- **Preprocessed data** (CSV files with suffix "\_preproc\_curves"): raw data that underwent the pre-processing procedure, such as median filtering or

shrinking (see [Analysis](#)). The curves will have blanks subtracted (if any specified in the layout) and samples marked as "Ignore" in the layout will be removed.

- **Fit curves** (CSV files with suffix "\_fit"): these curves are computed over the whole X-axis range of the input data using the determined fit parameters and are used to generate plots in the GUI and PNG format. This table is provided for convenience in case plotting of fit data outside MoltenProt is planned.
- **Baseline-corrected** (CSV files with suffix "\_raw\_corr"): raw data corrected for the pre- and post-transition baselines determined by the fit. These curves are useful for visualization and comparison between samples, because all Y-axis values are always in range from 0 (no protein unfolded) to 1 (all protein unfolded).

### 6.2.2. Fit parameters

The results of the fit are presented in two separate XLSX sheets/CSV files:

- **Fit parameters** (CSV files with suffix "\_results"): a table with all curve characteristics computed by MoltenProt. Parameters with suffix "\_init" are the initial parameters for the non-linear curve-fitting procedure; suffix "\_fit" marks the parameters obtained with curve fitting.
- **Standard deviations** (CSV files with suffix "\_results\_stdev"): non-linear curve-fitting procedure also yields a covariance matrix for obtained fit parameters. This information can be used to estimate the uncertainty of the fit and thus conclude if the current fitting result is reliable. For instance, well fit curves have Tm standard deviation of 0.5 K or below.

Initial values of the fit parameters have suffix "\_init", fit results have suffix "\_fit". Some parameters are shared by all built-in models of MoltenProt:

- **kN, bN**: slope and intercept of the pre-transition baseline, i.e. temperature dependence of native state (N) fluorescence.
- **kU, bU**: slope and intercept of the post-transition baseline, i.e. temperature dependence of unfolded state (U) fluorescence.
- **S**: standard error of the estimate (in units of Y-axis). This metric assesses how well the experimental data is described by the fit: for 99% of datapoints the difference between the observed and fit values will be below 3\*S. S is very sensitive for large outliers, such as spikes in the signal, so it should not be used as a single value to assess the curve quality
- **BS\_factor**: baseline separation factor (unit-less). BS-factor is a quantitative measure to assess how far is the pre-transition baseline from the post-transition baseline at Tm taking into account the noise in the signal (estimated via S). Curves with BS-factor above 0.8 are

exceptionally good, while curves with BS-factor in range 0-0.5 should be treated with caution. Negative BS-factor means that the curve is not suitable for interpretation. Since BS-factor is relative and unit-less, it is particularly helpful in deciding which readout to use for downstream analysis (e.g. F330 vs Ratio in NanoDSF data): the readout with higher average BS-factor is more preferable.

Model-specific parameters are described in [Models](#) section.

### 6.2.3. Sample information

The following sample information is included in the result table:

- **ID**: internal sample number assigned by MoltenProt. Up to 96 samples can be processed at once (A1 to H12).
- **Capillary** (only NanoDSF data): capillary position in the device.
- **Condition**: annotation describing the contents of the sample. For NanoDSF data the value is read from sheet "Overview". Annotations can be changed via [Layout editor](#).

## 7. Models

### 7.1. Overview

MoltenProt implements a variety of protein unfolding models, which should cover most common use-cases. All models rely on linear baseline extrapolation, which also means that data with sloping baselines should be processed with caution. This section contains a brief theoretical background for each model and introduces the recommended measure for final result ranking. All ranking measures are chosen in such a way that higher values correspond to higher stability of the protein.

### 7.2. Extensive and intensive readouts

Strictly speaking, the models implemented in MoltenProt are only applicable to extensive readouts, i.e. when the signal is proportional to protein concentration. While this is the case for raw fluorescence (330 or 350 nm) and Scattering, the F350/F330 Ratio is an intensive readout, because it is a proxy for the shape of the fluorescence spectrum. Applying equations for an extensive readout to an intensive readout produces an additional systematic error [4], [5]. The Ratio readout, however, tends to produce the most clean and easy to interpret sigmoidal curves, so omitting it from analysis decreases the explanatory power of the assay. For NanoDSF data MoltenProt also calculates deltaF readout (difference between fluorescence at 350 and 330 nm), which represents a trade-off between correctness and robustness. On the one hand, deltaF is an extensive readout, because it is a linear combination of F330 and

F350. On the other hand, subtraction of fluorescences removes a significant part of the baseline drift and can make the unfolding transition more pronounced.

## 7.3. Equilibrium models

Equilibrium models rely on several assumptions: 1) protein unfolding is a reversible reaction; 2) at every timepoint of the measurement the system is at chemical equilibrium; 3) protein heat capacity change ( $\Delta C_p$ ) is temperature-independent.

### 7.3.1. santoro1988

This model (applied to chemical denaturation) was initially described in ref. [6]. A more elaborate discussion for derivation of formulas is in ref. [5]. The model assumes that the protein exists in either native (N) or unfolded (U) state and there is an equilibrium between the folding and unfolding reactions ( $N \rightleftharpoons U$ ). The law of signal  $F(T)$  is described by equation:

$$F(T) = \frac{k_N \cdot T + b_N + (k_U \cdot T + b_U) \cdot \exp\left(\frac{\Delta H_m}{R} \cdot \left(\frac{1}{T_m} - \frac{1}{T}\right)\right)}{1 + \exp\left(\frac{\Delta H_m}{R} \cdot \left(\frac{1}{T_m} - \frac{1}{T}\right)\right)}$$

where  $k_N$ ,  $b_N$  are slope and intercept of the pre-transition (native) baseline,  $k_U$   $b_U$  are slope and intercept of post-transition baseline,  $R$  is the universal gas constant,  $\Delta H_m$  is enthalpy of unfolding at melting temperature  $T_m$ .

The final ranking metric is  $dG_{std}$ : Gibbs free energy of unfolding extrapolated to the standard state temperature (298.15 K).  $dG_{std}$  integrates the slope and the inflection point of an unfolding curve into a single measure.

By default extrapolation to the standard state temperature is linear, which is equal to the assumption that  $\Delta C_p$  is zero. If  $\Delta C_p$  is known, the extrapolated  $dG_{std}$  can be corrected by adding  $\Delta C_p * dCp\_component$ .  $dCp\_component$  is automatically calculated in santoro1988 mode and added to the output.

### 7.3.2. santoro1988i

This model is based on the same assumptions as the previous one, however, three states are possible for the protein: native (N), unfolded (U) and unfolding intermediate (I). If the  $T_m$  for N and I states is significantly different, it is possible to see two unfolding transitions in the experimental curve (two peaks on the smoothed derivative curve). Derivation is also described in ref [7].

The law of signal is as follows:

$$F(T) = \frac{k_N \cdot T + b_N + k_I \cdot \exp\left(\frac{\Delta H_{m1}}{R} \cdot \left(\frac{1}{T_1} - \frac{1}{T}\right)\right) + (k_U \cdot T + b_U) \cdot \exp\left(\frac{\Delta H_{m1}}{R} \cdot \left(\frac{1}{T_1} - \frac{1}{T}\right)\right) \cdot \exp\left(\frac{\Delta H_{m2}}{R} \cdot \left(\frac{1}{(T_1 + \Delta T_{2-1})} - \frac{1}{T}\right)\right)}{1 + \exp\left(\frac{\Delta H_{m1}}{R} \cdot \left(\frac{1}{T_1} - \frac{1}{T}\right)\right) \cdot \exp\left(\frac{\Delta H_{m2}}{R} \cdot \left(\frac{1}{(T_1 + \Delta T_{2-1})} - \frac{1}{T}\right)\right)}$$

where  $k_N$ ,  $b_N$  are slope and intercept of the pre-transition (native) baseline,  $k_U$   $b_U$  are slope and intercept of post-transition baseline,  $k_I$  is the signal slope for the I state (the state is assumed to be short-lived, so the intercept is not modelled),  $R$  is the universal gas constant,  $\Delta H_{m1}$  and  $\Delta H_{m2}$  are enthalpy of unfolding at melting temperature  $T_1$  and  $T_2$  (melting temperature for  $N \leftrightarrow I$  and  $I \leftrightarrow U$  reactions). In order to enforce the correct order of melting temperatures ( $T_2$  following  $T_1$ ),  $T_2$  is fit as  $T_1 + \Delta T_{2-1}$  (difference between  $T_2$  and  $T_1$ ).  $\Delta T_{2-1}$  is set to be non-negative.

The final ranking metric is `dG_comb_std`: Gibbs free energy of unfolding extrapolated to the standard state temperature (298.15 K), which is a sum of `dG_std` for reactions  $N \leftrightarrow I$  and  $I \leftrightarrow U$  (thermodynamic coupling).

## 7.4. Empirical models

Empirical models describe sigmoidal curves that are common in thermal unfolding assays without providing insights about the properties of the protein molecules.

### 7.4.1. santoro1988d

This model is a "descriptive" version of santoro1988 model: instead of enthalpy of unfolding at  $T_m$  ( $\Delta H_m$ ), the model uses onset temperature  $T_{onset}$  to describe the steepness of the curve.  $T_{onset}$  is the temperature at which 1% of protein is unfolded. The exponent in santoro1988 model is thus substituted to the following expression:

$$\exp\left(\frac{(T - T_m) \cdot \log\left(\frac{0.01}{0.99}\right)}{T_{onset} - T_m}\right)$$

The final ranking metric is the square root of the sum of squared  $T_m$  and  $T_{onset}$ . This can be thought of as the Euclidean distance from the point 0,0 K of a scatter plot between  $T_m$  and  $T_{onset}$ . The samples that are most far away from this point will have the most beneficial combination of  $T_m$  and  $T_{onset}$ . This ranking assumes that  $T_m$  and  $T_{onset}$  are equally important for protein stability.

See also ref. [8].

## 7.4.2. santoro1988di

This model is a "descriptive" version of santoro1988i model. The exponents with  $\Delta H_{m1-2}$  and  $T_{m1-2}$  are substituted to exponents using  $T_{onset1-2}$  and  $T_{m1-2}$  (see santoro1988d). Similarly to santoro1988i, this model can describe unfolding curves with two transitions.

The final ranking metric is the sum of geometric means of  $T_{m1}$  and  $T_{onset1}$  and  $T_{m2}$  and  $T_{onset2}$ .

## 7.5. Kinetic models

Kinetic models describe protein unfolding reactions via reaction rate constant, which links conversion of reactants to products with time. Arrhenius equation is used to model the temperature dependence of the reaction rate constant.

### 7.5.1. irrev

This model assumes that protein exists in states N and U only and the unfolding reaction is irreversible ( $N \rightarrow U$ ). The law of signal  $F(T)$  is defined as follows:

$$F(T) = k_U \cdot T + b_U + (k_N \cdot T + b_N) \cdot x_N(T)$$

where  $k_N$ ,  $b_N$  are slope and intercept of the pre-transition (native) baseline,  $k_U$   $b_U$  are slope and intercept of post-transition baseline and  $x_N(T)$  is fraction of natively folded molecules as a function of temperature.  $x_N(T)$  is obtained via numeric integration:

$$x_N(T) = \int_{T_{min}}^{T_{max}} \frac{-1}{v} \cdot \exp\left(\frac{-E_a}{R} \cdot \left(\frac{1}{T} - \frac{1}{T_f}\right)\right) \cdot x_N$$

where  $T_{min}$  and  $T_{max}$  are the start and end temperatures of the measurement,  $v$  is the scan rate (degrees/min),  $E_a$  is activation energy of unfolding,  $T_f$  is the temperature where reaction rate constant of unfolding ( $k$ ) equals 1,  $R$  is the universal gas constant.  $x_N$  is assumed to be 1 at  $T_{min}$ . See refs [7], [9] for derivation of similar equations for differential scanning calorimetry data.

The final ranking metric for this model is the negative logarithm of the reaction rate constant at standard state temperature ( $pk\_std$ ). Similarly to  $dG\_std$  in santoro1988, this metric integrates slope and inflection point of the unfolding curve (represented here as  $E_a$  and  $T_f$ ) in a single measure of stability.

### 7.5.2. lumry\_eyring

The Lumry-Eyring model [10] assumes that the protein exists in three states: native (N), unfolded (U) and aggregated (A). The reaction  $U \rightarrow A$  is irreversible. Two more non-equilibrium reactions  $N \rightarrow U$  and  $U \rightarrow N$  with reaction rate constants  $k_F$  and  $k_R$  describe the transition from state N to state U. Fitting the whole model to typical experimental data is not stable, so in MoltenProt this model is applied in two steps:

1. Scattering data (part of NanoDSF datasets if the respective detector is available) is fit with irrev model to obtain activation energy Ea and Tf (temperature where the rate constant is 1) for reaction U → A. It is assumed that states N and U produce the same Scattering signal.
2. The obtained parameters are supplied to the equation that describes protein unfolding signal F(T) (e.g. F350/F330 ratio in NanoDSF data) as a function of fraction native state xN, fraction unfolded state xU and fraction aggregated state xA.

$$F(T) = (k_N \cdot T + b_N) \cdot x_N + k_I \cdot x_U + (k_U \cdot T + b_U) \cdot x_A$$

$$x_N + x_U + x_A = 1$$

where kN, bN are slope and intercept of the pre-transition (native) baseline, kU bU are slope and intercept of the post-transition baseline (which is in fact represented by the A state, but not U), kI is the slope for the U state of unfolding. The equations for xU and xA are obtained with numeric integration [7].

The final ranking metric for this model is the negative logarithm of the ratio of the reaction rate constants for reactions N → U and U → N calculated at standard state temperature (pk\_ratio\_std). Similarly to dG\_std in santoro1988, this metric integrates slope and inflection point of the unfolding curve in a single measure of stability. The special feature of lumry\_eyring model in MoltenProt is that it can integrate not only the information from individual readouts, but also combine the stability as measured by Scattering with one selected unfolding readout.

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### 8.2. Dependencies

#### 8.2.1. pandas

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## 8.2.2. numpy

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