PyDSC Handbook

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1 Comments on the DSC Experiment

To ensure an optimal data correction and use of the script, the following procedure was applied during the experiment (see a summary Fig. 1).

- First of all, for the standard sample/buffer runs, the mass of solution in each DSC cell (sample and reference) was measured. From this information and the known concentration of our sample, the following masses should be known: in the sample cell; sample and buffer mass, in the reference cell; buffer mass. Attention must be paid to minimize the difference of mass between the buffer of each cell, $\Delta m_{BS} \sim 0$.
- Then, buffer/buffer runs were conducted. Same mass measurements as previously were applied, so the buffer mass is known in each cell. However, in that case, a mass difference of around 10% of the total volume must be ensured to have a clear signal, $\Delta m_{BB} \sim 0.1 m_S$.
- Finally empty cell/empty cell runs were performed. To have the most rigorous corrections possible, the empty cells should be the same all along these three runs, and each of them kept as sample or reference definition during all experiments.

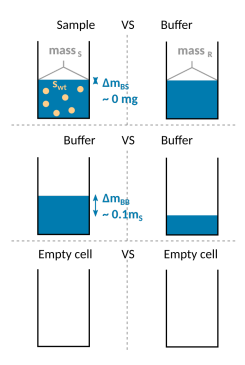


Figure 1: Scheme of the different runs in a DSC experiment. mass_S refers to the mass of the solution in the sample cell, mass_R to the mass of the buffer in the reference cell. s_{wt} is the concentration of the sample in weight percent. Δm_{BS} and Δm_{BB} are the difference of buffer mass between the sample and the reference cell, respectively in sample/buffer runs and buffer/buffer runs.

2 PyDSC

2.1 Theory

The script is written in Python 3, it makes use of the SciPy Package[1] and is platform independent. Several operations on the data are performed, provided the required information are available. The following operations are executed on the raw data files:

- 1. **Data averaging and resizing:** The DSC raw thermograms of the sample and, if provided, of the empty cell and buffer-buffer runs are read by the program and the points which fall outside the temperature region of interest are discarded. The reference measurements are averaged. If requested by the user, in order to facilitate any further data treatment, the number of points is reduced by a factor N, defined by the user.
- 2. **Heat flow conversion in heat capacity:** The heat flow is converted into a heat capacity upon division by the heat rate. The heat rate is either provided by the user, or it is determined from the raw data file if also the time information is contained. The raw data are finally plotted.
- 3. Empty cell and buffer correction: If the data file corresponding to the empty cell/empty cell and/or buffer/buffer runs are provided, the raw data are corrected for both contributions. In particular, the empty cell/empty cell run is used to take into account any non-constant contribution arising from the empty cells or the instrument. The buffer/buffer run is used to take into account any contribution arising from a mismatch in amount of buffer contained in the reference and in the sample cell. In detail, the apparent heat capacity of the sample run Cp_s , is corrected for the apparent heat capacity of the empty cell Cp_{EC} and for the apparent heat capacity of the buffer/buffer run Cp_{bb} as:

$$\overline{Cp_s} = Cp_s - Cp_{EC} - \frac{\Delta m_{BS}(Cp_{bb} - Cp_{EC})}{\Delta m_{BB}}$$
(1)

with Δm_{BS} and Δm_{BB} being the mass difference between the buffer contained in the sample and in the reference in the sample run, and between the buffers in the buffer/buffer run. If no empty cell/empty cell or buffer/buffer runs are provided, this correction will not be performed and the non-corrected sample run will be further analysed. In most of the cases, Δm_{BS} should be so small that the last term might be negligible.

- 4. Normalisation of apparent heat flow: The apparent heat flow of the sample run is finally normalized by the amount of solute. In particular, the apparent heat capacity is normalized either by the mass of solute, resulting in a Cp_s given in J K⁻¹ g⁻¹, or by the moles of solute, if the molecular weight is known, thus resulting in a Cp_s given in J K⁻¹ mol⁻¹.
- 5. Baseline correction: The core task carried out by the program is to compute a physically meaningful baseline. The baseline is constructed following an iterative procedure proposed elsewhere [2, 3]. In detail, the baseline $CP_{bl}(T)$ is defined as follows:

$$CP_{bl}(T) = (1 - \alpha)CP_{bl}^{pre}(T) + \alpha \cdot CP_{bl}^{post}(T)$$
(2)

with $CP_{bl}^{pre}(T)$ and $CP_{bl}^{post}(T)$ are the baselines determined in the regions at lower and higher temperatures than the peak, respectively. α is the degree of convertion, defined as:

$$\alpha(T) = \frac{\int_0^T CP_s(T) - CP_{bl}(T)dT}{\int_0^\infty CP_s(T) - CP_{bl}(T)dT}$$
(3)

where $CP_s(T)$ is the apparent heat capacity difference between sample and reference, after normalization and eventual correction. $CP_{bl}^{pre}(T)$ and $CP_{bl}^{post}(T)$ represent the temperature dependent difference of heat capacity of sample and reference, eventually corrected by the contribution from the empty cells and solvent mismatch. They are assumed to exhibit a linear dependence from temperature.

At this stage of the program, the enthalpy change of the process, computed as $\Delta H = \int_0^\infty C P_s(T) - C P_{bl}(T) \, dT$, is evaluated and its uncertainty is estimated. The uncertainty in ΔH is estimated from the stochastic fluctuations of the signal in the regions where $C P_{bl}^{pre}(T)$ and $C P_{bl}^{post}(T)$ are determined. By definition, in these regions no processes take place, and the signal fluctuates around an apparent heat capacity of zero J K^{-1} , which is used to determine the standard deviation of the heat capacity. The heat exchanged during the examined process is computed by integration using the trapezoidal rule, and its uncertainty by standard error propagation.

2.2 General use of the script

- 1. Save your thermograms in the associated folder rawdata.
- 2. Open Files.txt and enter the names of the files to analyze, separated by comma.
- 3. Open Input_params.txt and fill it according to your experiment (see the part 2.3 for more details).
- 4. Check that an Output folder is already created (it will be necessary to have this folder, so the script can save the results in it, or else it will return an error).
- 5. Run the script following one of the following procedures (with Anaconda Spyder or from a terminal).

2.2.1 From Anaconda - Spyder

- 6. Install Anaconda or Miniconda (see webpage¹ for the installation). The different libraries (numpy, scipy) and the editor Spyder will be automatically installed.
- 7. Run Spyder and open pyDSC.py
- 8. Run the file (play button in the top or press F5)

2.2.2 From a terminal

- 6. Install Python 3 and the associated libraries in your computer (or check if already done)
- 7. Open a terminal and go to the folder of interest with cd command
- 8. Write: python3 pyDSC.py
- 9. The script will be run, messages concerning the advancement will be displayed in the terminal

The output files will be saved in the Output folder and pdf files will be created for the graphs (see the part 2.4 for more details).

¹https://www.anaconda.com/distribution/

2.3 Input files

2.3.1 Files.txt

Fill with the names of the rawdata files, for heating and cooling cycles, and for each type of run.

- Sample heating cycles: Names of heating sample/buffer runs, separated by commas.
- Empty cell heating cycles: Names of heating empty cell/empty cell runs, separated by commas.
- Buffer heating cycles: Names of heating buffer/buffer runs, separated by commas.
- Sample cooling cycles: Names of cooling sample/buffer runs, separated by commas.
- Empty cell cooling cycles: Names of cooling empty cell/empty cell runs, separated by commas.
- Buffer cooling cycles: Names of cooling buffer/buffer runs, separated by commas.

Leave empty if no corresponding measurement was performed.

2.3.2 Input_params.txt

Fill with the parameters of the experiment.

Dataformat: refers to the format in which the dara are saved. It depends on which type of instrument is used. The options are:

Setaram3: raw ascii data which, in addition to the header of arbitrary length, contain three columns: Time(s), Furnace Temperature (°C or K), and HeatFlow (mW). The script will detect the «Furnace» word in the header to determine the first line of data. Example data are:

Sample name

Creation Date : 10/16/2018 4:33:29 PM

User : PSCM

HeatFlow :

Initial Mass: 456 mg

Molar Mass : N/A

Time (s)	Furnace Temperature (°C)	<pre>HeatFlow (mW)</pre>
0	4.843128	-0.006023
4	4.82847	-0.005714
8	4.691505	-0.004915
4792	85.045596	-0.522189
4796	85.114593	-0.520797
4800	85.183724	-0.521634

Setaram3temptime: raw ascii data, which, in addition to the header of arbitrary length, contain three columns: Furnace Temperature (°C or K), Time(s), and HeatFlow (mW).

The script will detect the «Furnace» word in the header to determine the first line of data. Example data are:

Sample_name

Creation Date: 10/16/2018 4:33:29 PM

User : PSCM

HeatFlow :

Initial Mass: 456 mg

Molar Mass : N/A

Furnace Temperature (°C)	Time (s)	<pre>HeatFlow (mW)</pre>		
4.843128	0	-0.006023		
4.82847	4	-0.005714		
4.691505	8	-0.004915		
• • •				
85.045596	4792	-0.522189		
85.114593	4796	-0.520797		
85.183724	4800	-0.521634		

Setaram4: raw ascii data, which in addition to the header of arbitrary length, contain four columns: Index, Time(s), Furnace Temperature (°C or K), and HeatFlow (mW). The script will detect the «Furnace» word in the header to determine the first line of data. Example data are:

Sample name

Creation Date : 10/16/2018 4:33:29 PM

User : PSCM

HeatFlow :

Initial Mass: 456 mg

Molar Mass : N/A

1	Time (s)	Furnace Temperature 4.843128	(°C)	-0.006023
2 3 	8	4.82847 4.691505		-0.005714 -0.004915
1199 1200	4792 4796	85.045596 85.114593		-0.522189 -0.520797
1201	4800	85.183724		-0.521634

3cols: raw ascii data with a **one-line header** which contain the data in a three column structure: Time(s), Furnace Temperature (°C or K), and HeatFlow (mW). In that case the script will consider by default a fixed header of one line, and will not look for «Furnace» word. It avoids error messages in the import process if you do not have a «Furnace» word in your header. Example data are:

Time (s)	Temperature	(°C)	${\tt HeatFlow}$	(mW)
0	4.843128	-0	.006023	
4	4.82847	-0.0	005714	
8	4.691505	-0.0	004915	
4792	85.045596	-0.	522189	
4796	85.114593	-0.	520797	
4800	85.183724	-0.	521634	

In all these cases, we assume that **the columns are separated by space or tab space**. Either your instrument is directly giving you .txt or .dat files (among others) in one of the Setaram format, either you need to format them by hand (for example, from an excel file, copy and paste the three columns of interest <u>in the right order</u> with a one-line header in an empty .txt or .dat file, and use **3cols** option).

- ROI_h: is the temperature region of interest for the heating runs. Fill with: Initial temperature, Final temperature, in °C. The data points from heating scans falling outside this area will be discarded.
- ROI_c: is the temperature region of interest for the cooling runs. Fill with: Initial temperature, Final temperature, in °C. Can be different from the heating runs. The data points from cooling scans falling outside this area will be discarded.
- mass_s: refers to the mass of the solution in the sample cell (see mass_s in Fig. 1), in mg.
- mass_r: refers to the mass of the buffer in the reference cell (see $mass_R$ in Fig. 1), in mg.
- mass_bb: refers to the difference of buffer mass Δm_{BB} in the buffer/buffer runs between the sample and reference cell (see Fig. 1), in mg.
 - s_wt: is the concentration of the sample in the sample/buffer run (see s_{wt} in Fig. 1). It is expressed in weight percent, so for example 1% wt will be 0.01, 100% wt will be 1.0.
- Scanrate_h: is the scan rate of the heating runs in K/min. If the data contain the time and temperature information, this line will be ignored and an average scan rate will be calculated from the time and temperature data.
- Scanrate_c: is the scan rate of the cooling runs in K/min. If the data contain the time and temperature information, this line will be ignored and an average scan rate will be calculated from the time and temperature data.
 - bins: represents the size of bins used to reduce the file size, i.e., a file of length N is reduced to N/bins, whereby bins number of points are averaged.
 - Encoding: Encoding used to decode the inputfile. Most common values are utf-8 and utf-16.
 - ROP_h: is the temperature region where the peak is found in the heating scans,. Outside this region, linear fits will be performed in order to evaluate the theoretical baseline. Before this region, it will be $CP_{bl}^{pre}(T)$, and after this region, $CP_{bl}^{post}(T)$. Fill with Initial temperature, Final temperature in °C.
 - ROP_c: is the temperature region where the peak is found in the cooling scans. Outside this region, linear fits will be performed in order to evaluate the theoretical baseline. Before this region, it will be $CP_{bl}^{pre}(T)$, and after this region, $CP_{bl}^{post}(T)$. Fill with Initial temperature, Final temperature in °C.

Mw: is the molecular weight of the sample in g/mol. It is optional.

Input: Convention used for input files, can be exo-up or exo-down.

Output: Convention used for input files, can be exo-up or exo-down.

Exo_in_plot : True or False, defines weather an arraw, indicating convention exo-up or exo-down are drawn on the graphs.

2.4 Output files

2.4.1 pdf files

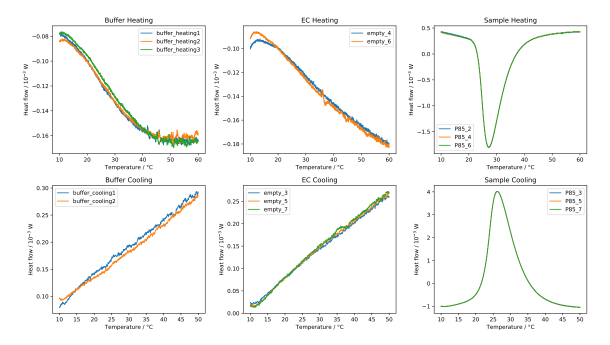


Figure 2: Rawdata.pdf shows the rawdata given by the instrument.

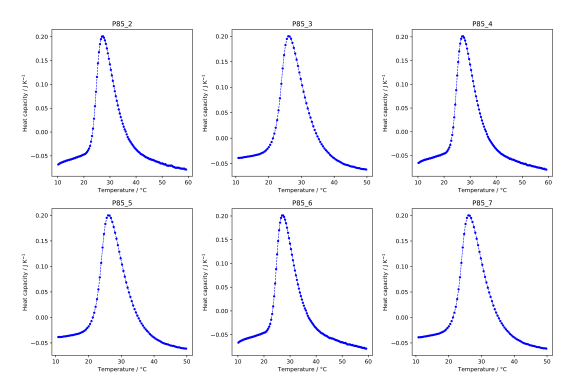


Figure 3: Corrected_data.pdf shows the data corrected by buffer and empty cell, and normalized by the scan rate (but not normalized yet by the sample mass). These curves help to visualizing the data points, and checking the binning.

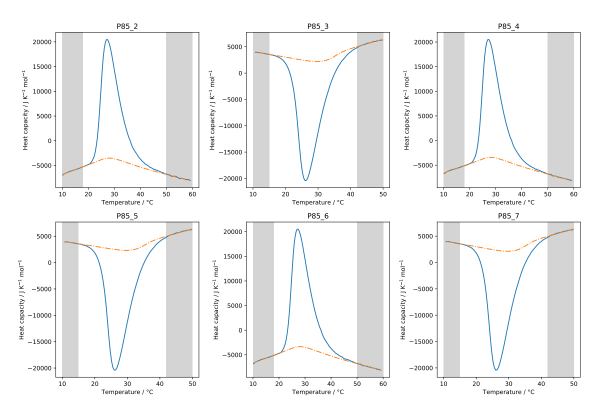


Figure 4: Baseline_data.pdf shows the corrected data, along with the baseline in orange dotted line, constructed following Eq. (2). The grey areas correspond to the respective range of pre-peak and post-peak linear fits used in Eq. (2). If there is no clear peak to create a baseline, like in the figure in the right, put a very small ROP (in Input_params.txt, for example here we chose «69.9, 70.0» for ROP_c), and the script will not try to create a baseline.

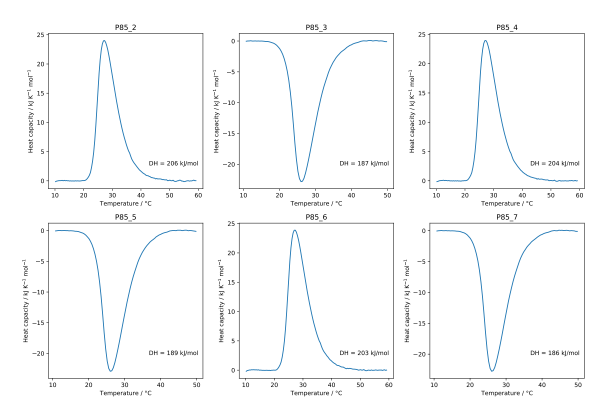


Figure 5: Final_data.pdf shows the thermograms after correction and after subtraction of the baseline. It is the data used for calculating the excess enthalpy ΔH , displayed in the figure.

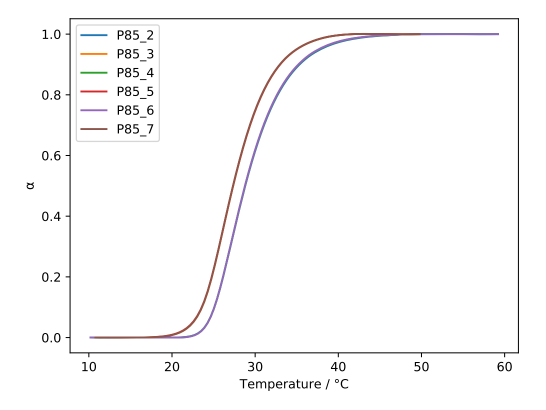


Figure 6: alpha.pdf shows the degree of conversion determined for each sample run, as a function of temperature.

2.4.2 txt files in the Output folder

For each experiment, the script will create a txt file displaying a header, explaining how the data were corrected and displaying thermodynamic parameters ΔH , T_m and ΔC_p , as well as five columns:

- 1. Temperature column Temp/ [degC], in °C.
- 2. The final heat capacity CP-baseline / [J/K/g] or CP-baseline / [J/K/mol], corresponding to the corrected data subtracted by the baseline (the ones displayed in Fig. 6), in J/K/g or in J/K/mol if the molecular weight is given in the input files.
- 3. The corrected heat capacity CP [J/K/g] or CP [J/K/mol], in J/K/g or J/K/mol (these data are shown in Baseline_data.pdf displayed in Fig. 4).
- 4. The final baseline <code>[J/K/g]</code> or <code>baseline [J/K/mol]</code>, in <code>J/K/g</code> or <code>J/K/mol</code>, constructed in an iterative way, following Eq. (2), and displayed in <code>Final_data.pdf</code>, shown in Fig. 4.
- 5. The enthalpy H [J/g] or H [J/mol], in J/g or J/mol, calculated as the integral over temperature of the final heat capacity (column 2): $H = \int_0^T [CP_s(T) CP_{bl}(T)] dT$.

If a molecular weight is added in the input file, it will display these data in J/mol.

3 Some functions of PyDSC (defined in DSC1.py)

3.1 correction

Corrects for empty cell VS empty cell, and buffer VS buffer runs following Eq. (1). More specifically:

- 1. It takes the average of all the scans performed for empty cell VS empty cell, and buffer VS buffer;
- 2. Then it does an interpolation of this average;
- 3. The correction as written in Eq. (1) is applied, taking the interpolations for empty cell and buffer contribution.

3.2 normalize_sampleruns

Normalizes the corrected heat flux by the sample mass (or eventually the molar mass if given in the input files), and the scan rate.

3.3 baseline

Creates the baseline in an iterative way as written in Eq. (2), calculates the enthalpy H, excess enthalpy ΔH and heat capacity difference ΔC_p (defined as the difference at transition temperature).

References

- [1] Eric Jones, Travis Oliphant, and Pearu Paterson. SciPy: Open source scientific tools for Python.
- [2] G Van der Plaats. A theoretical evaluation of a heat-flow differential scanning calorimeter. Thermochim. Acta, 12:77–82, 1984.
- [3] D V Malakhov and M. K. Abou Khatwa. Constructing a self-consistent integral baseline by using cubic splines. J. Therm. Anal. Calorim., 87(2):595–599, feb 2007.