Biophysical Characterization of Protein-Nucleic Acid Interactions

MicroScale Thermophoresis

Clemens Entzian, Corinna Kuttenberger, and Thomas Schubert

2bind GmbH, Regensburg
Our mission is to give researchers, developers, and decision makers the power to accelerate and streamline key research processes. We offer tailor-made biophysics services and solutions that are perfectly adjusted to your precise needs in the key, high-value areas of Drug Discovery, Antibody Research, Aptamer Characterization, and Protein Biophysics.

A state-of-the-art portfolio of biophysical technologies enables us to provide you with a complete and comprehensive picture of your specific question: From affinity and stability to kinetics and thermodynamics. This portfolio includes MicroScale Thermophoresis (MST), nano-Differential Scanning Fluorimetry (nanoDSF), Biolayer Interferometry (BLI), and Isothermal Titration Calorimetry (ITC).

Work with the Experts in Biophysics
MicroScale Thermophoresis (MST) is an optical fluorescent method. It records the changes in fluorescence of a target molecule as a function of temperature and the concentration of the cognate ligand molecule. Thus, in MST, the target must be fluorescent and the ligand must not be fluorescent in the same wavelength range as the target. The fluorescence changes are the result of two effects:

First, molecules exhibit directed movement along temperature gradients in solution, an effect called thermophoresis (Duhr & Braun, 2006). Such thermophoretic movement of a molecule is characterized by its size (hydrodynamic radius), its surface charge, and its hydration shell. All three parameters can be affected by binding of a ligand molecule. Thus, a concentration-dependent titration of the target molecule’s ligand induces changes in the thermophoretic movement. These changes translate in quantifiable, spatial fluorescence changes, which are easily tracked optically via the target fluorescence.

Second, fluorescence is a function of the temperature. Thus, the fluorescence of the target molecule varies with the temperature (Baaske et al., 2010). This temperature-dependence is additionally affected by the local molecular surroundings of the fluorophore. Thus, binding of a ligand molecule to the fluorescent target can change the chemical environment of the target fluorophore and thus the overall detected temperature-dependence of the fluorescence. This effect is known as TRIC (temperature-related intensity change; Gupta, Duhr & Baaske, 2018).

The thermophoresis and TRIC signals are additive and thus both contribute to the high sensitivity and robustness of MST measurements towards molecular binding events of all kinds. Thus, MST can be used for determining the affinity and binding strength of almost any kind of molecular interaction with very low sample consumption and very high sensitivity.

Figure 1. Technical MST setup. (A) MST measurements take place in small glass capillaries. Infrared and fluorescence lasers are used for generation of the MST effect and sample tracking. (B) TRIC and thermophoresis together account for a time-dependent change in fluorescence upon infrared-heating of the sample capillaries. (C) Multiple MST traces are recorded for different mixture ratios of target and ligand molecules. (D) Dose-response analysis of the MST traces allows for determination of the steady-state affinity of the target-ligand interaction.
**Features**

- Wide molecule size range
- Real-time quality controls
- No immobilization required
- Very short analysis time
- Free choice of assay buffer
- Low sample consumption
- Wide affinity range
- Wide temperature range

**Advantages**

- Analyze molecules from 100 Da up to 1 MDa
- Online aggregation, precipitation, and sticking controls
- Measurement is done truly in solution
- High analysis speed enables high throughput
- Biological liquids such as serum or cell lysate possible
- Minimum of only 6 µl per sample, nM concentrations
- Affinities can be analyzed in the pM-mM range
- Analysis possible from 20°C to 45°C

**Analysis of protein–nucleic acid interactions with MST**

Interactions of proteins and nucleic acids are reported to play a pivotal role in epigenetic networks, especially in the chromatin decondensation processes (Schubert et al., 2012). Dysregulated epigenetic networks represent a major reason for disease development and thus analysis of protein-nucleic acid interactions can deliver a multitude of new drug targets. It is often complicated to access the basic binding parameters such as steady-state affinity of nucleic acid fragments using biophysical methods.

Here, we demonstrate how MST can be used to determine the affinity of the epigenetic effector protein Df31 to single stranded (ss) RNA-, ssDNA-, or double stranded (ds) DNA fragments. The nucleic acid fragments shared the exact same sequence and length and differed only in their ss/ds “nature”.

For this, a simple yet robust MST assay was developed, in which the nucleic acid fragments were used as the constant target interaction partner, labeled with a Cy5 fluorescent dye for optical tracking during the measurements. When working with nucleic acids this labeling strategy often makes sense, because the nucleic acids can by directly synthesized with a fluorescent dye attached to their 5’ or 3’ ends. The Df31 protein was used as the ligand and titrated in 16 steps across a concentration range from 800 µM down to 350 nM.

We observed that Df31 specifically binds the ssRNA fragment with a $K_D$ value of 24 ± 3.1 µM and fails to recognize the single-stranded or double-stranded fragments (Figure 2). Importantly, the whole nucleic acid profiling just required single-digit nM concentrations of the nucleic acids. This can be of high importance when working with longer and expensive nucleic acids (e.g. with chemically modified bases). Although the used concentrations of the DF31 were quite high – in order to cover its affinity range with a full dose-response curve – the used volumes were very low. For each experiment, only 5 µL of the highest concentration solution were required. Thus, the whole profiling only required 15 µL of the Df31 protein.
Conclusion

In this application note, we demonstrated that our versatile MST assay is capable of measuring interactions between nucleic acids and binding proteins in a fast and robust manner as well as with minimal sample consumption. Similar assays can be readily developed for other questions of nucleic acid-protein binding.

<table>
<thead>
<tr>
<th>Assay type</th>
<th>MST</th>
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<tbody>
<tr>
<td>Target</td>
<td>Df31 ssRNA, Df31 ssDNA, Df31 dsDNA</td>
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<tr>
<td>Labeling</td>
<td>Cy5</td>
</tr>
<tr>
<td>Ligand</td>
<td>Df31</td>
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<tr>
<td>Measured $K_D$ [$\mu$M]</td>
<td>24 ± 3.1 (Df31 ssRNA) - (Df31 ssDNA) - (Df31 dsDNA)</td>
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Figure 2: MST dose-response curves of fluorescently labeled ssRNA, ssDNA, and dsDNA of the same sequence and length (39 nucleotides) to Df31. The ssRNA-Df31 dose-response curve was fitted to the Hill equation, data represent the mean ± standard deviation from three technical replicates.

Literature


