ABSTRACT

The Protein Thermal Shift Assay (TSA) is a rapid and sensitive tool for monitoring protein thermostability, aiding in the identification of optimal conditions or conformations/sequences that favour protein stability, including the concentration of protein-ligand interactions. TSA is based on temperature-induced protein denaturation, monitored using an environmentally sensitive dye, such as SYPRO® Orange. The fluorescence data is plotted and the midpoint or Tm of the resulting curve is taken as a reference of the thermal stability of a protein of interest. Comparisons can then be made between Tm values obtained using a range of buffer conditions or addition of different ligands.

TSA data have been obtained from the whole range of Applied Biosystems Real Time PCR Systems, including the 7900 HT, 7500 Fast™ and StepOnePlus™ Real Time PCR Instruments, demonstrating the versatility of these systems. The benefits of performing a TSA with an AB Real Time PCR System include the flexibility of run- method programs, catering for a range of data resolution requirements, and in the use of small reaction volumes, providing fast and accurate results with only a few µg of protein.

INTRODUCTION

The difficulty and expense often involved in isolating or obtaining a protein of interest, demands that care be taken with subsequent handling or storage in order to maximise the utility and longevity of the protein and to ensure data quality is not affected by degradation or aggregation events. Conditions that favour long-term stability are a common requirement for almost all research or applied techniques involving proteins. There are many factors that may affect protein stability, including salt concentrations, pH or the use of specific ligands that can interact with proteins in different ways. Given the large number of possible combinations one could test in order to determine the environmental conditions that would favour maximum stability, it is highly desirable to adopt a technique that can simplify and streamline the investigation process.

The Thermal Stability Assay (TSA) on AB Real Time PCR Systems is a fast and convenient screening method, and the use of a non-specific dye such as SYPRO® Orange alleviates any requirement for prior knowledge of protein function or ligand activity. In the presence of a native protein, the dye is naturally quenched, when the protein of interest starts to denature in response to an increase in temperature, exposing the hydrophobic core of the protein, the dye will react to this change in environment and will start to fluoresce (see Figure 1).

Figure 1. Schematic of the predicted action of SYPRO® Orange Dye. The naturally quenched dye will only fluoresce when exposed to the hydrophobic regions of a denaturing protein.

Fluorescence data is collected by the Real Time PCR instrument throughout the temperature ramp, a melt curve is generated, and the midpoint or melt peak can be determined for each well across a 96 or 384 well plate (Figure 2).

Comparisons of the resulting Tm values can be used to determine the relative stabilising properties of the test conditions.

Figure 2. Example TSA data from a StepOnePlus™ showing the Normalized Reporter and Derivative Melt profiles from experimental Lysosome samples. Data was collected at 1°C intervals from 25°C through to 99°C.

Figure 3. Replicate TSA data from a 96-well plate using the 7900 HT Real Time PCR System, demonstrating high level of reproducibility.

It is not necessary to calibrate an AB Real Time PCR System for the SYPRO Orange Dye, however calibration may improve results in some instances. Recommendations for calibration involve the use of a full plate of replicate conditions with a high enough concentration of protein to achieve maximum fluorescence (as described in the Maintenance Guides for each specific system), for example, with the 7900 HT Real Time PCR System, an rfu value of 30,000 would be sufficient.

TSA EXAMPLE EXPERIMENTS

The following examples were generated using an AB 7900 HT Real Time PCR instrument, using the Dissociation Curve run method and modifying the ramp rate as required.

Effect of pH on Protein Stability

The delicate tertiary structure of many proteins can be difficult to maintain during handling and storage. The pH of a chosen buffer will certainly affect protein stability as changes in pH alter the electrostatic interactions between charged amino acids. Using the screening approach of the TSA it is possible to test a range of pH conditions in a single assay. The changes in the resulting Tm of the protein in solution can be used as a measure of the relative stability across the tested pH range.

Figure 4a shows the normalized melt curve data generated from a single 96-well plate on the AB 7900 Real Time PCR System, containing duplicate reactions of a protein in solution across a pH range of 4.20 – 9.49. The Tm values of the TSA melt curves (omitting the two obvious outlier groups) span across a ten degree window, clearly demonstrating the effect of pH on the resulting thermal stability of the protein being tested. Figure 4b plots the Tm values obtained against the solution pH, where it can be determined that a buffer solution at pH 5.47 leads to the most stable conditions.

Figure 5. Derivative Melt Curve data from a 7900 HT Real Time PCR System depicting a TSA assay measuring the effect of salt concentrations on protein thermostability. The Tm of the protein of interest increases with different concentrations of salt in the buffer.

Effect of Ligand Interaction on Protein Stability

Ligands are small molecules or substances that may exhibit specific affinities for some proteins, and through the formation of the protein-ligand complex, can affect the stability of a protein of interest, for example by changing the protein structure or conformational stability. Ligands are often used to increase the success rate of protein purification or crystallization by increasing the stability of the protein being studied. Ligand-protein interactions are also highly relevant in drug screening procedures. There are a large range of ligands that may be of interest, and TSA is an attractive screening tool for measuring protein-ligand interactions, as there are no requirements to monitor enzyme activity, or even to have prior knowledge of the ligands being tested.

Figure 6 depicts triplicate reactions of a specific protein-ligand TSA. The normalized melt curves on the left of the graph have a Tm of around 55°C, representing native protein in a simple buffer solution containing no ligand. The three curves to the right represent the same buffer protein in the presence of a specific ligand, with Tm values around 10°C higher than the native protein alone, a clear demonstration of the increased stability afforded by this protein-ligand interaction.

Figure 6. Normalized Melt Curve data generated on a 7900 HT Real Time PCR System showing the effect of a specific ligand on protein thermostability. The Tm of the curve increases over 10°C in the presence of the ligand.

CONCLUSIONS

TSA is a rapid, inexpensive and straightforward tool for screening of suitable conditions that maximise protein thermostability. The use of AB Real Time PCR Instruments offers a range of benefits to assist the TSA testing workflow, combining the ease of handling and throughput capacity of a plate-based system with the ability to hold samples at sub-ambient temperatures. In addition, the flexibility of the operational software allows for the collection of fluorescent data across a wide temperature range at any desired ramp speed, with data easily viewed within the system software, or exported for further analysis offline.

References

1 SYPRO Orange Protein Gel Stain SKG85-S6550
2 Nisaii et al., - Published online 13 September 2007; doi: 10.1038/jmor.2007.312
3 Mrakowski et Protein Sci. 2006 15: 2712-2718

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