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Thermodynamics of Natural and Synthetic Inhibitor Binding to Human Hsp90

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1. Introduction

Heat shock protein 90 (Hsp90) is one of conserved heat shock proteins that protect, prevent aggregation, stabilize, activate, or otherwise regulate client proteins, is a component of the cellular chaperone machinery [Taipale et al., 2010, Taldone et al., 2009, Wandinger et al., 2008]. There are a number of recent developments in the understanding of the interesting and complex mechanism of Hsp90 action [Neckers et al., 2009a, Neckers et al., 2009b, Mayer et al., 2009, Walerych et al., 2009]. Hsp90 is over-expressed in cancer cells and Hsp90 inhibitors have shown selectivity for cancer cells. Therefore, small-molecule inhibitors are being developed as anticancer therapeutics [van Montfort and Workman, 2009, Sharp et al., 2007, Sgobba and Rastelli, 2009, Fukuyo et al., 2009].

Two groups of Hsp90 inhibitors have been designed based on naturally occuring inhibitors geldanamycin and radicicol. Geldanamycin has been modified to 17-AAG, while various resorcinol-bearing compounds were designed based on radicicol. Here we describe the thermodynamics of their binding to Hsp90 by isothermal titration calorimetry (ITC) and thermal shift assay (TSA). These assays yield not only the potency, i.e. the Gibbs free energy of binding, but also the enthalpy of binding, the entropy of binding, and the heat capacity of binding. This detailed thermodynamic description and the comparison between homologous compound structures, coupled with structural information of the Hsp90-inhibitor complex, provides insight into the structure-activity relationships (SAR) of the compounds. The SAR helps in the process of rational drug design [Freire, 2009].

2. The structure of Hsp90 and the comparison of human and yeast isoforms

There are several Hsp90 homologs in human, yeast, and bacteria. Human Hsp90 exists in two highly homologous isoforms - α and β. Alpha isoform is prevalent. There are no major known functional differences between the isoforms. Hsp90 homolog in yeast is named Hsc82 and also shares significant homology with human isoforms.

Figure 1 shows the structure of Hsp90 and Hsc82. The protein in solution exists in equilibrium between dimer and monomer. Furthermore, the protein is quite flexible and exists in equilibrium between at least three major conformations [Graf et al., 2009]. The full length protein consists of three major domains – the N-terminal domain (1-216 a.a.), the M-domain (262-524), and the C-terminal domain (525-709). There is also a charged linker (216-262) that did not crystalize and its structure is unknown [Ali et al., 2006]. Inhibitors
such as radicicol and geldanamycin bind to the ATP-binding pocket of Hsp90 and are thus competitive non-covalent inhibitors.

Fig. 1. The structure of Hsp90. Top left. The superimposition of the N-termini of human Hsp90αN (orange) and Hsc82N (blue) with transparent surfaces show that the fold is essentially identical. Radicicol is shown bound to yeast Hsc82N (PDB ID: 1bgq) as spacefilled green model. Top right. Rotation of the ‘lid’ part of the protein (94-139 a.a., 108-139 a.a. in human Hsp90). Bottom left. The structure of Hsc82 full length dimer (2cg9). One monomer is shown in dark blue-red and another in light blue-pink. Blue shows the position of the N-terminal domains. Bottom right. The structure of Hsc82 full length monomer showing the positions of three domains and the unstructured charged linker.
Radicicol interactions with human and yeast Hsp90 isoforms are shown in Figure 2. Crystallographic experimental structure exists only for the yeast isoform. Therefore, radicicol binding to human isoforms was modelled computationally into the active site.

Fig. 2. Interactions of radicicol (grey substance in the middle at the top of the picture) with selected Hsp90 amino acids. Hsp90α amino acids are shown in orange (3eko) superimposed on yeast Hsc82 amino acids (colored by atom, 1bgq), and Hsp90β amino acids are shown in pink. Three important water molecules participating in hydrogen bond formation between the protein and radicicol are shown as spheres.

There are very few amino acid differences between yeast and human isoforms that may have an impact on the binding thermodynamics. They are shown in the figure and listed below:

Hsp90α  Ser52  Ile91  Val92  Ala141  His154
Hsp90β  Ala47  Leu86  Val87  Ala136  His149
Hsc82  Ala38  Ile77  Arg78  Leu127  Ser140

All other amino acids in the vicinity of radicicol are identical in all three isoforms. Therefore, the difference in the binding thermodynamics should be due to the above mentioned amino acids. Furthermore, charged amino acids Arg78 and His154 point to the solvent on the
proteins surface different from radicicol binding site. Therefore, the difference is most likely
due to the remaining three amino acids.

3. Thermodynamics of binding by isothermal titration calorimetry and
thermal shift assay

Protein – ligand binding equilibrium is described by the Gibbs free energy of binding \( \Delta_G \).
More negative \( \Delta_G \) indicates a stronger binding reaction. The Gibbs free energy is sufficient
to describe the equilibrium. However, several thermodynamic parameters that contribute to
the \( \Delta_G \) can be correlated with structural features of the protein – ligand complex easier
than the \( \Delta_G \) itself. The most important parameters are the enthalpy \( \Delta_H \) and entropy
\( \Delta_S \) of binding:

\[
\Delta_G = \Delta_H - T \Delta_S
\]  

Both the enthalpy and entropy are the first temperature derivatives (T-derivatives) of the
Gibbs free energy:

\[
\left( \frac{\partial \Delta_G}{\partial T} \right)_P = -\Delta_S
\]  

\[
\left( \frac{\partial \ln K}{\partial \frac{1}{T}} \right)_P = \frac{\Delta_H}{R}
\]

The second T-derivative of the \( \Delta_G \) (the \( \Delta_H \) T-derivative) is the heat capacity of binding
\( \Delta_C_p \). Subscript P indicates constant pressure.

\[
\frac{\partial \Delta_H}{\partial T} \right)_P
\]

There are other thermodynamic parameters that are pressure derivatives (P-derivatives) of
\( \Delta_G \). The first P-derivative is the volume of binding \( \Delta_V \). The second P-derivatives are
the compressibility and expansion of binding. These parameters may be measured by
varying the pressure of the protein – ligand system. However, they are rarely used and are
beyond the scope of this chapter. Here we will concentrate on the most used, however,
selected thermodynamic parameters, namely, \( \Delta_G, \Delta_H, \Delta_S, \) and \( \Delta_C_p \).

The Gibbs free energy of ligand binding may be measured by a large variety of methods,
well reviewed for carbonic anhydrase inhibitor binding in [Krishnamurthy et al., 2008]. Here
we will concentrate on the application of ITC and the (TSA). Both methods have been
described previously in detail, especially ITC [Freyer and Lewis, 2008, Landbury, 2004,
ITC directly measures the heat evolved or absorbed during the binding reaction. At constant pressure, the heat is equal to the enthalpy ($\Delta_h H$) of binding. This method is the most robust and accurate way of measuring the $\Delta_h H$. However, until the isothermal titration calorimeters became commercially available in early 90s, the $\Delta_h G$ was usually estimated from the $\Delta_h G$ T-dependence using the van’t Hoff relationship (3). If all contributing reactions are clearly dissected, such approach should yield the same results as titration calorimetry. However, in practice, there are many unexplained inconsistencies and only ITC provides reliable $\Delta_h H$.

However, the ITC has a number of disadvantages. Most importantly, the binding constant should be in a rather narrow range to satisfy the requirement that coefficient $c$ is between about 5 and 500. The $c$ is:

$$c = nM_t K_b$$  \hspace{1cm} (5)

Where $n$ is the binding stoichiometry, $M_t$ is the protein molar concentration, and $K_b$ is the binding constant defined for the reaction of $M + L \rightleftharpoons ML$ as:

$$K_b = \frac{[ML]}{[L][M]}$$  \hspace{1cm} (6)

$$\Delta G = -RT\ln K_b$$  \hspace{1cm} (7)

In practice, ITC is useful for $K_b$s in the range of $10^5$ to $10^9$ M$^{-1}$. Such $K_b$s can be usually measured by varying protein concentration. If ITC experiment is planned well, it can provide $\Delta_G$, $\Delta_h H$, and $\Delta_S$ in an hour. Doing the same experiment at several temperatures will yield an indirect measurement of the heat capacity $\Delta_C_p$.

Another disadvantage of ITC is that it requires rather large amount of protein (usually more than 0.1 mg) and ligand. The protein must be well purified and soluble at micromolar concentrations.

These disadvantages can be quite easily approached using the TSA. This method is based on the observation that specifically binding ligands stabilize (sometimes destabilize) the protein. Protein solution is being heated at a constant rate in the absence or presence of a ligand and the unfolding pattern is measured by various methods such as absorbance, circular dichroism, or, most often, by fluorescence. Various fluorescent components could be followed, such as intrinsic tryptophan fluorescence or an extrinsic solvatochromic probe. Most convenient is 1,8-anilino naphthalene sulfonate (ANS). Figure 3 shows ANS fluorescence dependence on temperature upon Hsp90 unfolding. The rise in fluorescence near 50 °C is due to Hsp90 unfolding and the exposure of hydrophobic patches of the protein interior. ANS binds to such patches and its fluorescence increases.
Fig. 3. Thermal shift assay protein melting curves. The midpoint of denaturation ($T_m$) is shifted to higher temperature upon ligand addition.

Note that ANS primarily binds to cationic groups on the protein surface (first to arginine residues) [Matulis and Lovrien, 1998]. However, most such bound ANS does not fluoresce and thus we can observe the unfolding pattern of the protein. Addition of ligand stabilized the protein and shifted the curve and the midpoint of the unfolding transition ($T_m$) by about 5 degrees.

Protein unfolding fluorescence curves are described by the equation:

$$y = y_N + \frac{y_U - y_N}{e^{\Delta\frac{RT}{y}} + 1} + \frac{y_N - y_U}{e^\frac{RT}{y}}$$  \hfill (8)

Protein melting temperatures can be determined by fitting the protein melting curves (Figure 3) according to:

$$y(T) = y_{T_r} + m_f(T - T_m) + \frac{(y_{T_r,T_m} - y_{T_r,T_m}) + (m_{r} - m_f)(T - T_m)}{1 + e^{(\frac{y_{T_r,T_m} - y_{T_r,T_m}}{RT}) + \frac{\Delta m}{RT}}$$  \hfill (9)
where \( y(T) \) is the calculated fluorescence as a function of temperature; \( y_{f,T_m} \) is the fluorescence of the probe bound to folded native protein before the transition at \( T_m \); \( y_{u,T_m} \) is the fluorescence of the probe bound to the unfolded protein after the unfolding transition at \( T_m \); \( m_F \) is the slope of the fluorescence dependence on temperature when the probe is bound to the native protein; \( m_U \) is the slope of the fluorescence dependence on temperature when the probe is bound to the unfolded protein; \( \Delta_U H_{T_m} \) is the enthalpy of protein unfolding at \( T_m \); \( \Delta_U S_{T_m} \) is the entropy of protein unfolding at \( T_m \); \( \Delta_U C_p \) is the heat capacity of protein unfolding and is assumed to be temperature-independent over the temperature range studied; \( R \) is the universal gas constant; and \( T \) is the absolute temperature (Kelvin).

Ligand dosing curves (as in Figures 9 and 10) are described by the equation (10):

\[
L_T = \left( \frac{K_{u,T_m}}{2K_{u,T_m}} \right) \left( \frac{P_T}{K_{b,T_m}} + \frac{1}{K_{b,T_m}} \right) \left[ \frac{1}{e^{-\Delta_U H_{T_m} + \Delta_U C_p(T_m-T_r) - RT_m(\Delta_U S_{T_m} + \Delta_U C_p \ln(T_m/T_r))} - 1} \right]
\]

\[
\times \left[ \frac{P_T}{2} e^{-\Delta_U H_{T_m} + \Delta_U C_p(T_m-T_r) - RT_m(\Delta_U S_{T_m} + \Delta_U C_p \ln(T_m/T_r))} + \frac{1}{e^{-\Delta_U H_{T_m} + \Delta_U C_p(T_m-T_r) - RT_m(\Delta_U S_{T_m} + \Delta_U C_p \ln(T_m/T_r))} - 1} \right]
\]

\[ L_T \] is the total concentration of added ligand, \( K_{u,T_m} \) is the protein unfolding equilibrium constant at \( T_m \); \( P_T \) is the total protein concentration; \( K_{b,T_m} \) is the ligand binding constant at \( T_m \); \( \Delta_U H_{T_m} \) is the enthalpy of protein unfolding at \( T_m \); \( T_r \) is the protein melting temperature when no ligand is added; \( \Delta_U S_{T_m} \) is the entropy of protein unfolding at \( T_m \); \( \Delta_U C_p \) is the heat capacity of protein unfolding and is assumed to be temperature-independent over the temperature range studied; \( \Delta_U H_{T_0} \) is the enthalpy of ligand binding at \( T_0 \); \( T_0 \) is the temperature at which the binding process is studied (usually 37 °C); \( \Delta_U S_{T_0} \) is the entropy of ligand binding at \( T_0 \); \( \Delta_U C_p \) is the heat capacity of ligand binding and is assumed to be temperature-independent over the temperature range studied.

The binding constant at the physiological temperature \( T_0 \) is determined using:

\[
K_{b,T_0} = e^{\frac{\Delta_U H_{T_0} - \Delta_U S_{T_0} \ln(T_0/T_r)}{RT_0}}
\]

TSA can be performed in the RT-PCR machine and requires only several micrograms of protein. Furthermore, there is no upper limit of the \( K_b \) to be determined. The only limit is the temperature of water boiling. Therefore, such extremely tight reactions as radicicol binding to Hsp90 can be studied by TSA. There is also no lower limit for the \( K_b \). Therefore, millimolar and picomolar ligands can be easily measured. However, TSA does not determine \( \Delta_U H \), \( \Delta_U S \), and \( \Delta_U C_p \). Therefore, both TSA and ITC should be used to determine the thermodynamics of Hsp90 – ligand binding.

4. Thermodynamics of Hsp90 ligand binding

Figure 4 shows the structures of Hsp90 ligands used in this study.
Fig. 4. Chemical structures of compounds discussed in this manuscript
Fig. 5. A. Isothermal titration calorimetry data for radicicol binding to Hsp90αN. Upper graph – raw ITC data, lower graph – integrated ITC data with the curve fit to the standard single binding site model. The cell contained 4 μM protein, while the syringe contained 40 μM radicicol in the same buffer - 50 mM sodium phosphate, pH 7.5, 0.5% DMSO, 100 mM NaCl, at 25 °C. B. Radicicol binding ITC curve at the same conditions as in panel A except pH 8.5. C. 17-AAG binding to Hsp90αN. D. ICPD47 compound binding to Hsp90αN
Fig. 6. The observed enthalpies as a function of the buffer deprotonation enthalpy at 25 °C temperature in various buffers: △ – phosphate, ● – Hepes, ▲ – Tris. The data points are the experimentally-observed enthalpies, and the trendlines are linear fits. Their slopes are equal to the binding-linked protonation events. Intersection with y axis is buffer-independent binding enthalpy.

Fig. 7. Intrinsic binding enthalpies obtained after accounting for the linked protonation event as a function of temperature for radicicol binding to Hsp90αN (■) and Hsp90αF (○). The slopes are linear fits to the experimental data and are equal to the intrinsic heat capacities of radicicol binding.
Figure 8. The binding of AZ3 to Hsp90αN at pH 7.0 as determined by the thermal shift assay. Upper panel shows experimental fluorescence curves. The lower panel shows the same curves recalculated as probabilities to observe the protein in the denatured state.

Figure 5 shows several typical ITC binding curves of the Hsp90 – ligand system. The curve in panel A is too steep, meaning that radicicol binding is too tight to be accurately determined by ITC. TSA data will be needed to determine the $K_b$ and $\Delta G$ of interaction. However the $\Delta H$ is determined to high precision. The $\Delta H$ determined in buffers with different enthalpy of protonation ($\Delta H_a$) yielded different $\Delta H$ (Figure 6). Therefore, the binding reaction is linked with protein or ligand protonation or deprotonation upon binding. In other words, ligand binding shifts the $pK_a$ of ionisable groups as previously explained [Baker and Murphy, 1996].
Due the linked protonation, it is important to dissect protonation thermodynamics from binding thermodynamics in order to determine the intrinsic thermodynamics of binding. The enthalpy of, for example, Tris buffer protonation is so large (about -44 kJ/mol) that it would hide any binding enthalpies. Therefore, a series of experiments in various buffers are necessary (Figure 6).

When protonation-linkage effects are accounted for [Zubriene et al., 2010] and the intrinsic enthalpy of binding is determined, such experiments should be repeated at all temperatures of interest. Figure 7 shows intrinsic enthalpies determined at 13, 25, and 37 °C for radicicol-Hsp90α system. The full length protein bound radicicol with slightly less exothermic enthalpy. The difference was equal to ~4 kJ/mol (Table 1). This difference is within the standard error of the measurements. The error was greater for the full length protein because the available protein amount and concentration was lower.

ITC was useful to provide the enthalpy of binding. However, as seen in radicicol binding ITC curves in Figure 5, the binding is too tight and would require the displacement assay as described previously [Velazquez-Campoy and Freire, 2006]. Our experience shows that the TSA is much easier and yields more precise results than the displacement ITC assay [Zubriene et al., 2009].

Therefore, the binding of all ligands listed in Figure 4, was measured by the TSA. Figure 8 shows typical raw protein melting curves observed at various added ligand concentration. AZ3 bound with relatively low, millimolar affinity. Therefore, relatively large concentration of ligand had to be added in order to observe the $T_m$ shift. It should be noted here that the shift continues way beyond saturation of protein with ligand. This is due to the dominant entropy of mixing. If the stabilization occurred due to some kind of bond formation, then we would not observe continued stabilization past the saturation point. This is observed when an inhibitor binds covalently and irreversibly to the protein.

Figures 9 and 10 show the dosing curves for the ligands listed in Figure 4. Radicicol, the most potent binder, shifts the temperature by nearly 15 °C. 17-AAG is the average binder and shifts the temperature by about 10 °C. Some ligands, such as AZ1, barely shift the temperature even at 1 mM concentration.

| Protein    | $K_d$, nM | $\Delta G_{int}^i$, kJ/mol | $\Delta H_{int}^i$, kJ/mol | $\Delta S_{int}^i$, J/K mol$^{-1}$ | $\Delta C_p$, J/K mol$^{-1}$ |
|------------|-----------|-----------------------------|-----------------------------|-----------------------------------|-----------------------------|
| Hsp90αN    | 0.04      | -70.7                       | -59.4                       | -11.4                             | -38                         |
| Hsp90αF    | 0.04      | -66.8                       | -59.4                       | -7.5                              | -25                         |
| Hsp90βN    | 0.15      | -60.6                       | -56.1                       | -4.6                              | -15                         |
| Hsc82F     | 0.25      | -46.7                       | -54.8                       | 8.1                               | 27.1                        |
| Uncertainties | ±1.6-fold | ±4                          | ±2.6                        | ±4.7                              | ±16                         |

Table 1. Intrinsic thermodynamic parameters of radicicol binding to human and yeast isoforms of Hsp90.
Fig. 9. The dependence of the melting temperature of Hsp90αN on the concentration of various inhibitors (ligand dosing curves) [Ugele et al., 2009]. The observed \( K_d \) (µM) by TSA were: Radicicol – 0.00083, AZ3 – 5000, ent-35 – 0.27, and 35 – 0.4

Fig. 10. The dependence of the melting temperature of Hsp90αN on the concentration of inhibitors 17-AAG and ICPD47. The observed \( K_d \) (µM) by TSA were: 17-AAG – 0.3 and ICPD47 – 0.002
Figure 11 shows an interesting phenomenon observed in TSA when the concentration of ligand is lower than protein. When the concentration of ligand is insufficient to saturate all protein binding sites, the denaturation transition splits into two transitions – the first transition is due to free protein and the second is due to the liganded protein. Relative magnitudes of both transitions are proportional to the concentrations of free and liganded protein concentrations. This phenomenon has been also observed by DSC with weakly-binding ligands at relatively high protein concentrations [Sharke and Ross, 1988].

TSA enabled determination of sub-nanomolar binding potency of naturally occurring radicicol and strongly or weakly-binding synthetic compounds where ITC does not work. However, no dissection of proton linkage was done for ligands where ITC was not feasible. Therefore, only observed $K_d$s are obtained for such ligands without other thermodynamic information. After dissecting the proton linkage by ITC, it was shown that radicicol binds about 4 times more strongly to recombinant human Hsp90 alpha than to Hsp90 beta isoform. This reduction in affinity is caused primarily by less favorable enthalpic rather than entropic contributions. About 90% of the binding energy comes from the favorable enthalpic contribution and small opposing entropic contribution at physiological temperature (Table 1).

Detailed proton linkage and temperature analysis had to be performed to dissect buffer and protein linked reactions from ligand binding intrinsic reaction. However, even after this...
detailed analysis, it is not possible to determine whether conformational change in the protein could contribute significantly to these intrinsic thermodynamic parameters. It is quite likely that some contribution comes from the rotation of the lid as shown in Figure 1. The intrinsic enthalpy of radicicol binding to Hsp90 is one of the largest enthalpies observed for any protein - small ligand binding. Note, that most of the Gibbs free energy of radicicol binding comes from the favourable enthalpic contribution. The entropy contribution is relatively small.

5. Conclusions

Radicicol and other resorcinol-bearing compound binding to Hsp90 is interesting in many respects regarding drug design. First, the binding reaction can be very tight (i.e., it has a very favorable Gibbs free energy). Radicicol stabilizes Hsp90 by 15-20°C. Second, the binding reaction has a very favorable enthalpy of binding, one of the largest for any protein - small ligand system. Third, there are few direct contacts between Hsp90 and radicicol that could account for such a large binding energy. Fourth, water molecules play an essential role in the recognition and binding. And fifth, the negative heat capacity of binding usually reflects a dominant hydrophobic origin of binding. However, hydrogen bonds are apparently essential for radicicol binding to Hsp90.

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