

Thermodynamics (Classical) for Biological Systems

Prof. G. K. Suraishkumar

Department of Biotechnology

Indian Institute Of Technology Madras

Module No. # 06

Reaction Equilibria

Lecture No. # 34

Criteria for Bio-reaction Equilibria

Welcome back!

In this lecture, we will begin the last module for this course. The last module is module six on reaction equilibria. If you recall in the last module, we had explicitly said that we will not consider any reactions. ... We had situations like that in biological systems, plenty, and therefore we could use whatever we developed in that module to explain many situations. But as you all know reactions take place all the time. Whether it is macro-biological system, for example, a bio-process – there is reactions taking place. Or even at the cell level - the cell is alive and kicking only because of the thousands of reactions that take place in a cell.

Under ... reasonable conditions, we could still consider this cell as a continuum, and therefore apply these principles of analysis to the cell also. Therefore, the reactions that take place in the cell can also be considered to be a valid system for the application of these principles. So, let us start looking at the bases to develop reaction equilibria, the conditions for the same, or the criteria for the same. Earlier, if you recall, in module five, we had criteria for the phase equilibria or when there were no reactions that we are taking place, and so on. We will develop similar criteria, when reactions are present.

(Refer Slide Time: 02:06)


Let us consider a bio-chemical reaction, say that occurs during the cell metabolism, that is represented by the equation:

$$2 A + B \rightarrow 3 C \quad \text{Eq. 6.1}$$

The stoichiometry represented by the above equation can be represented as:

$$0 = 3C - 2A - 1B$$

or, in general as


$$0 = v_1 M_1 + v_2 M_2 + v_3 M_3 \quad \text{Eq. 6.2}$$

To do that, we will touch up on many things that you may already be somewhat familiar with, because of exposure in eleventh, twelfth standards, or even in courses that you took before ... you took this thermodynamics course. ... Let us consider a bio-chemical reaction, say that occurs during a cell metabolism, say a reaction in glycolysis. Let us represent that ... this is a hypothetical equation ... let us represent that by 2 A plus B giving you 3 C. I had just picked random numbers here, 2 A plus B giving you 3 C. Let us call that equation 6.1. The stoichiometry represented by the above equation can be written as 0 equals 3 C, 3 C minus 2 A minus 1 B. I have deliberately written it like this.

I have taken 3 to be positive, you know the product associated coefficient to be positive, minus 2 A minus 1 B equals 0. Or, in general, I could write this stoichiometry as, or I could represent the reaction as 0 equals sum nu 1 M 1 plus nu 2 M 2 plus nu 3 M 3. In a minute, I will tell you what nu s and M s and so on, are. In this case, nu 1 M 1, nu 2 M 2 nu 3 M 3. Let us call this equation 6.2, I wrote it ... in this form for a reason.

(Refer Slide Time: 04:21)

In other words

$$0 = \sum_i \nu_i M_i \quad \text{Eq. 6.3}$$


ν_i : the stoichiometric coefficient

+ ve for the products (the biochemical species on the RHS)
- ve for the reactants (the biochemical species on the LHS)

$$\nu_1 = -2, \nu_2 = -1, \nu_3 = 3$$

M_i : the biochemical species

For an inert species that is present in the system, but does not react, the stoichiometric coefficient is taken as zero – this will help in generalizations



I can now make this into a compact notation, 0 equals some over i nu i M i. This is the reason why I had written at that way; equation 6.3, where nu i is this stoichiometric coefficient which we will take as positive for the products of the reaction or the biochemical species on the right hand side of the equation. Like this; you know, if this is the equation, ... the coefficients on the right hand side, we are going to take as positive. And the coefficients on the left hand side, which is that for the reactance, we are going to take as negative.

So that works well here, 3 C was positive minus 2 A, reactant, negative, minus 1 B, reactant, negative. Therefore, negative for the reactants, which are the bio-chemical species on the left hand side. I have already given you an answer, but why did not you take about five minutes and complete it? Why did not you write down nu 1, nu 2 and nu 3 for the reaction that we just saw? Go ahead, take about five minutes, and write this down. Go ahead please.

(No audio from 05:39 to 10:59)

If you had followed the reasoning that we presented earlier, you can easily see that nu 1 was the coefficient of A, minus 2, nu 2, the coefficient of B, minus 1, and nu 3, the coefficient of C 3. And M i is the various biochemical species A, B, or C in this particular example that is considered in the reaction. Let me state this although we would not be looking at this in great detail, but it is good to state this here. If there is an inert species

that is present in the system, for an inert species, that is present in the system, but does not react, the stoichiometric coefficient is taken as zero. Have this in mind. Whenever it becomes necessary to use this, please use this, and that will make things a lot more general.

(Refer Slide Time: 12:07)


We know that the changes in the number of moles of each species are directly proportional to the corresponding stoichiometric coefficients

For example, in the above reaction, Eq. 6.1.

$$2 A + B \rightarrow 3 C \quad \text{Eq. 6.1}$$

if 1 mole of A disappears, 0.5 moles of B will also disappear to result in 1.5 moles of C

This can be represented as



Now, let us go back to what we already know. We know that the changes in the number of moles of each species in a reaction are directly proportional to the corresponding stoichiometric coefficients. What do I mean by that?

For example, in the above reaction given by equation 6.1, which is 2 A plus 3 B equals 3 C, if we divide this equation throughout by 2, we get A plus half B equals three by two C. Or, if 1 mole of A disappears, half a mole of B will also disappear to result in three by two or 1.5 moles of C. Therefore, the changes in the number of moles of each species are directly proportional to the corresponding stoichiometric coefficients.


(Refer Slide Time: 13:16)

$$\frac{\Delta n_1}{\nu_1} = \frac{\Delta n_2}{\nu_2} = \frac{\Delta n_3}{\nu_3} = \dots = \frac{\Delta n_i}{\nu_i} = \dots = \Delta \varepsilon \quad \text{Eq. 6.4}$$

Δn_i : the change in the number of moles of a species, i
 $\Delta \varepsilon$: the extent of reaction, or the reaction coordinate

For a differential change in the number of moles

$$\frac{dn_1}{\nu_1} = \frac{dn_2}{\nu_2} = \frac{dn_3}{\nu_3} = \dots = \frac{dn_i}{\nu_i} = \dots = d\varepsilon \quad \text{Eq. 6.5}$$

 NPTEL

This particular aspect can be represented as the change in number of moles of 1, Δn_1 , divided by the stoichiometric co-efficient, ν_1 , equals the change in the number of moles of 2 divided by the stoichiometric coefficient 2, equals the change in the number of moles in 3 divided by the stoichiometric coefficient 3 and so on, equals Δn_i divided by ν_i , and let us say that equals something called $\Delta \varepsilon$. We will call this equation 6.4.

Δn_i , as I said, was the change in the number of moles of a species i , and ... $\Delta \varepsilon$ is the extent of a reaction. It is also called the reaction coordinate, if you recall this term from your earlier courses. $\Delta \varepsilon$ - the reaction coordinate, or ε is the reaction coordinate.

For a differential change in the number of moles, we can replace these big differences by the differential change. And therefore, we can write dn_1 by ν_1 equals dn_2 by ν_2 , equals dn_3 by ν_3 and so on, which equals dn_i by ν_i , and that equals $d\varepsilon$. We will call this equation 6.5. ...What I would like you to do is now, take some time, I will tell you how much time. Look at glycolysis, the various reactions in glycolysis, which is one of the central pathways in the cell,

Remember ... glucose goes to glucose six phosphate. Glucose from outside the cell gets inside the cell first and then it will go to glucose six phosphate, fructose six phosphate and so on, all the way down to pyruvate. Choose that pathway, which is called

glycolysis. Each one of those steps as you know is catalyzed by an enzyme. Take those individual reactions and write equivalent expressions for, let us say, five of those reactions. I am going to give you about twenty minutes to do that. Take about twenty minutes; go back to your biochemistry text book, biochemistry notes. Choose five reactions that take place in glycolysis and write this down, for improving the comfort level in using this particular equation. Go ahead please.

(No audio from 16:14 to 36:39)

Now, hopefully, you would have gotten back the level of comfort that you had in your earlier classes while dealing with stoichiometric coefficients, and so on, by writing down the relationships between the stoichiometric coefficients for at least five reactions in glycolysis. The choice of the reactions were yours. Of course, they were reversible reactions but that does not matter.

(Refer Slide Time: 37:17)

Let us consider a system in which only the reaction taking place is Eq. 6.1

The Gibbs free energy of the reacting system is

$$G^T = n_1\mu_1 + n_2\mu_2 + n_3\mu_3 \quad \text{Eq. 6.6}$$

We know from the earlier courses on basic Thermodynamics that the equilibrium condition corresponds to a minimum in the Gibbs free energy

Since the composition (number of moles of each species) varies as the reaction proceeds, we can think of a certain composition at which the Gibbs free energy is a minimum

NPTEL

Now, let us look at the condition for bio-reaction equilibrium. Let us consider a system in which the only reaction taking place is equation 6.1, which is two A plus B giving you three C; that was the reaction given by 6.1. The Gibbs free energy of the reacting system is G^T as we had seen earlier. That can be written as $n_1\mu_1$, the chemical potential, you know, this is per unit mole μ_1 . Therefore, $n_1\mu_1$ will give you the total free energy corresponding to species 1; plus $n_2\mu_2$, the total free energy

corresponding to species 2, plus $n_3 \mu_3$, the total free energy corresponding to species 3, or species A, species B, and species C. Let us call this equation 6.6.

Now, we know from either chemistry courses, the thermodynamic aspects of chemistry courses, or some specific courses that you would have taken earlier that the equilibrium condition corresponds to a minimum in the Gibbs free energy. I am just going to take this directly from whatever we know earlier that a minimum in Gibbs free energy occurs at equilibrium. Since, the composition or the number of moles of each species varies as the reaction proceeds, we can think of a certain composition at which the Gibbs free energy is a minimum.

You know the reaction is going to take place, when the reaction takes place, with time, the number of moles of each species is going to change. We already know from earlier that equilibrium occurs when the Gibbs free energy is a minimum. In other words, there must be some combination of moles of these species involved in the reaction at which the Gibbs free energy turns out to be a minimum. That is the whole basis of the argument here.

(Refer Slide Time: 39:50)

This means that if G^T is plotted versus, say n_1 , at constant temperature and pressure, a minimum in G^T occurs at some value of n_1 , and that corresponds to the equilibrium condition

In other words, at equilibrium

$$\left(\frac{\partial G^T}{\partial n_1}\right)_{T,P} = 0 \quad \text{Eq. 6.7}$$

NPTEL

This means that if G^T , the total free energy, is plotted as a function of let us say n_1 , the number of moles of species 1 – that is the only thing that we can represent on one coordinate. That is only reason for taking n_1 . At constant temperature and pressure, a minimum in G^T occurs at some value of n_1 , and that corresponds to the equilibrium

condition. If we write it mathematically, at equilibrium, dn_i or dn_1 of G_T , total Gibbs free energy at constant temperature and pressure equals 0. This is just a mathematical statement of the fact that we knew from earlier classes that the free energy, Gibbs free energy, of a system is minimum at equilibrium.

Therefore, the slope of the G_T versus dn_1 curve goes to zero at that particular point. You know it is a minimum and therefore, this slope goes zero. Let us call this equation 6.7.

(Refer Slide Time: 41:06)

Further, we know from Eq. 2.15 that

$$dG^T = -S^T dT + V^T dP + \sum_i \mu_i dn_i \quad \text{Eq. 2.15}$$

Writing Eq. 2.15 for our current system of interest, we get

$$dG^T = -S^T dT + V^T dP + \mu_1 dn_1 + \mu_2 dn_2 + \mu_3 dn_3 \quad \text{Eq. 6.8}$$

NPTEL

Further, we know from equation 2.15 that dG_T equals minus $S_T dT$ plus $V_T dP$ plus sum over i $\mu_i dn_i$; you recall this equation? This equation we have used many times. If we write equation 2.15 for our current system of interest, which is the system that consists of that only reaction taking place, we can write dG_T ; the first two terms are, of course, minus $S_T dT$ plus $V_T dP$; and sum over i $\mu_i dn_i$ is plus $\mu_1 dn_1$ plus $\mu_2 dn_2$ and plus $\mu_3 dn_3$. Let us call this equation 6.8.

(Refer Slide Time: 42:12)


Applying Eq. 6.5

$$\frac{dn_1}{v_1} = \frac{dn_2}{v_2} = \frac{dn_3}{v_3} = \dots = \frac{dn_i}{v_i} = \dots = d\varepsilon \quad \text{Eq. 6.5}$$

to our current system, we get

$$\frac{dn_1}{-2} = \frac{dn_2}{-1} = \frac{dn_3}{3} \quad \text{Eq. 6.9}$$

And thus,

$$dG^T = -S^T dT + V^T dP + \left(\mu_1 + \mu_2 \left(\frac{-1}{-2} \right) + \mu_3 \left(\frac{3}{-2} \right) \right) dn_1 \quad \text{Eq. 6.10}$$


Now, if we apply 6.5, equation 6.5, which is dn_1 by ν_1 equals dn_2 by ν_2 equals dn_3 by ν_3 and so on, ... and it equals dn_i by ν_i – and we said we could equate it to $d\varepsilon$ of a reaction co-ordinate, $d\varepsilon$ to our current system – we get dn_1 by minus 2, and minus 2 was the stoichiometric coefficient of A; dn_2 by minus 1, 1 was the stoichiometric coefficient of B. And, since it is a reactant, this becomes minus one. Similarly, here also 2 was a stoichiometric co-efficient of A, and since it is a reactant, by our convention, this becomes minus 2; equals dn_3 by this stoichiometric coefficient of the product C, and since it is a product we have a positive terminology here, plus 3. So, dn_1 by minus 2 equals dn_2 by minus 1 equals dn_3 by 3.

Let us call this equation 6.9. ... Therefore, dG^T can be written as, minus $S^T dT$ plus $V^T dP$ plus this $\mu_1 dn_1$ plus $\mu_2 dn_2$ plus $\mu_3 dn_3$ can be written as $\mu_1 dn_1$ is fine. $\mu_2 dn_2$ is nothing but minus 1 by minus 2 dn_1 . That is what we have written here, minus 1 by minus 2 dn_1 . There is a bracket here, and we have taken dn_1 common out. And $\mu_3 dn_3$ is nothing but in terms of dn_1 , 3 by minus 2 dn_1 , and that is what we have written here. Let us call this equation 6 10.

(Refer Slide Time: 44:27)

At constant temperature and pressure, G^T can be zero only if the last term is zero, or,

$$\mu_1 + \frac{1}{2}\mu_2 = \frac{3}{2}\mu_3$$


In other words,

$$0 = 3\mu_3 - 2\mu_1 - \mu_2 \quad \text{Eq. 6.11}$$

which can be compared, term by term, with the bioreaction stoichiometry:

$$0 = 3C - 2A - 1B$$

except that the species are replaced by their chemical potentials.



And at constant temperature and pressure, G^T can be zero. It can be zero, only if the last term is zero, because at constant temperature and pressure as dT is zero, dP is zero anyway at constant temperature and pressure. Therefore dG^T can be zero, only if this term goes to zero. And therefore, $\mu_1 + \frac{1}{2}\mu_2$ must equal $\frac{3}{2}\mu_3$ or in other words, 0 must equal $3\mu_3 - 2\mu_1 - \mu_2$. We will call this equation 6.11, which can be compared, term by term with the bio-reactions stoichiometry as 0 equals $3C - 2A - 1B$, except that, this species are replaced by their chemical potentials. Can you see this here? $3C - 2A - 1B$, whereas here for the condition of equilibrium, we got $3\mu_3 - 2\mu_1 - \mu_2$.

(Refer Slide Time: 45:52)

In general, if there are k species in a bioreaction, written as

$$0 = \sum_i \nu_i M_i$$


then

$$dG^T = -S^T dT + V^T dP + \mu_1 dn_1 + \mu_2 dn_2 + \dots + \mu_k dn_k$$

which can be written in terms of the reaction coordinate and stoichiometric coefficients, from Eq. 6.5 as

$$dG^T = -S^T dT + V^T dP + (\mu_1 \nu_1 + \mu_2 \nu_2 + \dots + \mu_k \nu_k) d\varepsilon$$

Eq. 6.12



Therefore, in general, if there are k species in a bio-reaction, written as $0 = \sum_{i=1}^k \nu_i M_i$. This sum is taken from i equals 1 to k $\nu_i M_i$. Then, dG^T equals minus $S^T dT$ plus $V^T dP$ plus $\mu_1 dn_1$ plus $\mu_2 dn_2$ plus so on, till $\mu_k dn_k$. This is where we said that there are k species in the bio-reaction. If you can write this in terms of the reaction co-ordinate and the stoichiometric coefficients, from equation 6.5, we get dG^T equals minus $S^T dT$ plus $V^T dP$ plus $\mu_1 \dots$ you know this, dn_1 we could write as $\nu_1 d\varepsilon$; dn_2 we could as $\nu_2 d\varepsilon$, and so on. Therefore, $\mu_1 \nu_1 + \mu_2 \nu_2$ and so on plus $\mu_k \nu_k$ times $d\varepsilon$.

I hope you are able to see this. Go back to equation 6.5, if you are unable to see this, this will drop out from there. The relationship between the various dn and $d\varepsilon$ will drop out from there. And from that, we will get this expression. We will call this equation 6.12.


(Refer Slide Time: 47:29)

Therefore, at constant T and P , we can write

$$\left(\frac{\partial G}{\partial \epsilon}\right)_{T,P} = \sum_i \nu_i \mu_i \quad \text{Eq. 6.13}$$

G^T minimum occurs at equilibrium
For a G^T minima, the derivative in Eq. 6.13 should be zero
Thus, the condition for equilibrium is

$$\sum_i \nu_i \mu_i = 0 \quad \text{Eq. 6.14}$$

 This turns out to be the most fundamental condition to be satisfied for a reaction to be at equilibrium

Therefore, at constant temperature and pressure, we can write $\left(\frac{\partial G}{\partial \epsilon}\right)_{T,P}$ at constant T and P of G equals $\sum_i \nu_i \mu_i$. That is equation 6.13. We know that G^T minimum occurs at equilibrium and for a minima to occur the ... derivative given here must be equal to zero. And therefore, the condition for equilibrium is that $\sum_i \nu_i \mu_i$ equals 0, equation 6.14. And this happens to be the most fundamental equation to represent chemical equilibrium, or equilibrium of a system in which chemical reactions take place. So, please remember this. I think we are almost out of time.


(Refer Slide Time: 48:40)

If there are R independent reactions that occur in a system

For each of the independent reactions, there will be an equation of the form given in Eq. 6.14

$$\sum_i \nu_i \mu_i = 0 \quad \text{Eq. 6.14}$$

Thus, R such reactions will define that system at equilibrium



Therefore, when we come back, in the next class or let me see whether we can go further a little bit. If there are R independent reactions that occur in a system, for each of those independent reactions, there will be an equation of the form 6. 14. There is $\sum \nu_i \mu_i$ equals zero. Thus, R such reactions will define that system at equilibrium. Therefore, we will have, for a multi- reaction system, also the condition for equilibrium remains the same. But we will have R such equations that specify the equilibrium. We will stop here and when we start the next class, we will continue with other aspects of reaction equilibrium.