# Metabolic Control Systems

This lecture will cover the basics of metabolic control, including how enzymes and transport systems control the fate of a metabolite and how systems are integrated to maintain homeostasis. These concepts should be a review of material covered in your Biochemistry classes. For further details please refer to the books on reserve for this course<sup>1,2</sup>

#### **Contents**

Learning Objectives Key Terms and Concepts 2 Control of Metabolic Flux 2 Cellular Transport Systems Types of Membrane Transporters 3 Powering Active Transport Enzymes 4 Thermodynamics 4 **Enzyme Kinetics** 5 Cofactors Integrated Control of Metabolism 6 Post-Translational Modification 7 Transcriptional Regulation

- <sup>1</sup> J M Berg, J L Tymoczko, and L Stryer. *Biochemistry*, volume New York. 2013. ISBN 0-7167-3051-0
- <sup>2</sup> Denise Ferrier. *Lippincott Illustrated Reviews: Biochemistry*. LWW, 1496344499, 7th edition, 2017. ISBN 1496344499

## Learning Objectives

- Explain what a rate limiting enzyme is, what a committed enzyme step is and what a reversible reaction is.
- Predict the differences in speed and persistence of allosteric, posttranslational and transcriptional regulation of metabolism.
- Describe the role of cellular transport in macromolecular regulation. Understand the differences between active and passive transport.

# Key Terms and Concepts

- Activation Energy
- Allosteric Regulation
- Cofactor
- Concentration Gradient
- Enzyme
- Rate Limiting Enzyme
- Transcription Factor
- Transporters (Active and Passive)

#### Control of Metabolic Flux

Cells need to control the rates at which nutrients are taken up, stored, or used and there are several ways by which this occurs. Here we will review the biochemistry of both nutrient transport and enzyme function. Understanding these concepts will be very important to understanding how the metabolic pathways we will discuss are controlled.

#### Cellular Transport Systems

First we will describe the ways in which cells control nutrient permeability. Most of the nutrients we will discuss<sup>3</sup> are unable to pass through the plasma membrane of the cell. Allowing or denying access to a nutrient is one way by which cells can control nutrient metabolism. Without these transport mechanisms we would be unable to absorb digested food, or transport nutrients from cell to cell. While we normally think of transporters as getting nutrients into or

<sup>&</sup>lt;sup>3</sup> The exceptions are sterols and some other lipids

out of a cell, they are also important within cells, for example getting pyruvate into the mitochondria, or storing calcium in the endoplasmic reticulum.

#### *Types of Membrane Transporters*

Membrane transporters are generally fairly specific for the molecule they transport. For example GLUT<sub>4</sub> transports glucose, but GLUT<sub>5</sub> transports fructose. Transporters can broadly be separated into two major types, passive transporters and active transporters. These can be differentiated by considering whether they work with or against the concentration gradient, with active transporters typically working against the concentration gradient.

Passive transporters allow for nutrients to pass down a concentration gradient into the cell. As an example, the liver expresses a glucose transporter named GLUT2. Glucose can either enter the liver (if there is more glucose in the blood than the liver) or exit the liver (if the reverse is true). Passive transporters will only allow a nutrient to enter a cell if there is less of the nutrient in a cell than in the blood. This is quite efficient for disposing of excess nutrients, such as after a meal, but is not effective in storing things away against a concentration gradient. It may seem like passive transporters are not regulated, but as we will see in the case of GLUT4, the amount of transporters at the cell surface can be controlled by cell signaling<sup>4</sup>. The rate of a passive transport is defined by three things, the gradient of the transported molecule, the number of transporters at the relevant membrane, and the efficiency of the transporter.

ACTIVE TRANSPORTERS can force nutrients into a cell against the concentration gradient. These transporters function like pumps and have to use energy of one sort or another to "power" the molecule into the cell. You may think that this is a bad idea, but there are lots of examples where this matters physiologically. One example is retaining salt. If your kidneys werent actively retrieving sodim out of urine and back into the blood, then you would rapidly lose osmotic pressure in your blood. The key is to think about the concentration inside or outside, if you are pushing against the transport gradient, you need active transport.

#### Powering Active Transport

Active transport requires energy of some type. This energy can come from several sources such as ATP, other concentration gradients, or even light. Some examples are described in Figure 1. The key

4 if you want to jump ahead, here is a review on that process [Leto and Saltiel, to controlling the rate of these transporters is not only the concentration gradient of the transported molecule, but also the levels (or gradient) of the powering force. In the cases where molecules are co-transported they can either be pulled in simultaneously (this is known as a symporter) with the molecule of interest as shown on the left of Figure 1, or can be exchanged where one molecule exits, powering the entry of the molecule of interest (this is known as an antiporter). A classic example of an antiporter is the sodium:glucose exchanger SGLT1, which extrudes sodium down its concentration gradient (into the gut lumen) to force uptake of glucose from the gut into cells. This allows for efficient carbohydrate uptake in a meal<sup>5</sup> even if the gastrointestinal cells have similar or higher glucose levels to the gut lumen.

#### **Enzymes**

#### **Thermodynamics**

EVERY CHEMICAL IN THE BODY HAS A CERTAIN AMOUNT OF ENERGY. When we eat, some of this chemical energy is converted into ATP to allow for function. This is known as catabolism. When we are storing nutrients, we use ATP to generate higher energy molecules such as fats or glycogen. This is known as anabolism. Every molecule in our body has a set amount of energy and a chemical reaction can be considered endothermic (requiring energy) or exothermic (releasing energy), depending on whether the reactants or products have higher energy. The levels of these metabolites at equillibrium can be calculated with the following equation where  $K_{eq}$  is the *Eqiullibrium constant*<sup>6</sup>:

$$K_{eq} = \frac{[B]}{[A]} \tag{1}$$

The equillibrium constant can be calculated from the *free energy* of the reactants and products.

$$\Delta G_o = G_o' - RT ln K_{eq} \tag{2}$$

$$\Delta G_{o}^{'} = G_{o}^{'}(reactants) - G_{o}^{'}(products) \tag{3}$$

Some reactions have products with very similar energy levels and the balance between the reactants and the products is based primarily on their concentrations. This is known as an *equillibrium* reaction which would have a  $K_{eq}$  of near to 1. If a reaction requires a lot of energy to occur, this is often an *irreversible* or *committed* step<sup>7</sup>. This

<sup>5</sup> SGLT2 does a similar thing, retrieving glucose from urine back in to the blood. Therefore inhibiting SGLT2 prevents glucose retrieval back to the blood, and is the target of several drugs which try to lower blood glucose in diabetics. The trade names for these drugs include Invokana, Farxiga and Jardiance.

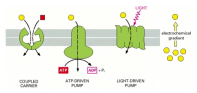


Figure 1: Examples of active transport. Reproduced from Alberts et al. [2002]

<sup>&</sup>lt;sup>6</sup> The square brackets mean the concentration of A or B

<sup>&</sup>lt;sup>7</sup> This would have a large, negative K<sub>eq</sub>

means that once this reaction happens, there is no going back. If you think about the metabolic pathway in Figure 2, this would mean that once you proceed through step 2 to make C you cannot go back to B. Given the free energy  $(G_0)$  and concentration of the reactants and products in a reaction you can calculate the  $\Delta G$  and equillibrium constant for a reaction and estimate whether it is reversible<sup>8</sup> under normal conditions.

#### Enzyme Kinetics

Without enzymes, many reactions occur very slowly due to the activation energy needed for the reaction to occur. Enzymes increase the rate of a chemical reaction by reducing the activation energy required for a reaction to occur. This does not change the equillibrium constant, it just allows the reaction to reach equillibrium faster. This is sketched out in Figure 3, note that  $\Delta G$  is not changed, but the dashed line has a higher activation energy, and therefore slower reaction rate than the solid line.

Most metabolic pathways are controlled by altering the RATES at which metabolites are converted to its final product. The overall rate of a metabolic pathway is controlled by the rate-limiting step<sup>9</sup>. In a linear pathway, the speed of this step's enzymatic reaction controls the overall rate. Quite often the rate-limiting enzyme is an important point of regulation, as adjusting the speed of this reaction can speed up or slow down and entire pathway.

Reaction rates increase in rate as the concentration of substrates increase until the enzymes are saturated (see the solid line in Figure 4). This is known as Michaelis-Menten kinetics. The reaction rate constant (k) and rate can be calculated from the activation energy with these equations<sup>10</sup>:

$$k = Ae^{-\frac{E_a}{RT}} \tag{4}$$

$$rate = k \frac{[Reactant][Enzyme]}{[Reactant] + K_m}$$
 (5)

If products build up the reaction becomes more complex and now looks like this where  $K_p$  is the binding constant for the product:

$$rate = k \frac{[Reactant][Enzyme]}{[Reactant] + K_m \left\{ 1 + \frac{[Product]}{Kp} \right\}}$$
(6)

Allosteric regulation is another way by which enzymes CAN CONTROL REACTION RATES. Allosteric enzymes are generally multi-subunit enzymes that change their Km as more products bind. 8 There is a good blog post explaining how the steady state  $\Delta G$  is determined on this basis at http: //sandwalk.blogspot.com/2007/10/ aldolase-reaction-and-steady-state.

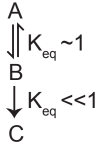


Figure 2: Example schematic of a metabolic pathway.

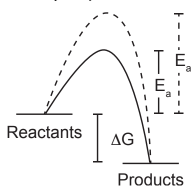


Figure 3: Example schematic of the activation energy (Ea) of an enzymatic reaction.

- 9 the rate-limiting step is generally the slowest step of a pathway. As an example, in glycolysis the rate-limiting step is catalysed by phosphofructokinase-1
- 10 A and R are constants, T is temperature and e is Euler's number.  $K_m$  is the Michaelis constant for an enzyme.

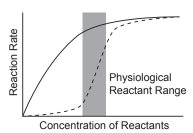


Figure 4: Example of Michaelis-Menten (solid line) and allosteric (dashed line) kinetics.

An example of this is the dashed line in Figure 4. This has several advantages in terms of regulation. One advantage is that the reaction rate can be effectively zero or at maximum in a much narrower range, bracketing the actual range of substrates present physiologically. Another advantage is that allosteric activators or inhibitors can shift the curve to the left or right, to effectively increase or decrease the reaction rate. This is a common mechanism by which that activity of rate-limiting enzymes are regulated.

On the basis of these equations, reaction rates (and the rate of a particular metabolic pathway) can be increased by several things<sup>11</sup>. Try to convince yourselves how this happens based on the equations listed above. Can you think of any other things that would affect pathway flux?

WHILE LINEAR FLOW THROUGH A PATHWAY IS IMPORTANT, another aspect of pathway control is how the fate of a particular nutrient is decided. This is illustrated in Figure 5. In the example on top the nutrient would be equally distributed between three products, but in the bottom example, by adjusting the rates of the specific pathways, a nutrient can be directed to a particular product. At several points during this class, we will describe how the "fates" of particular metabolites are controlled by the relative rates of metabolic pathways.

#### **Cofactors**

Many enzymes require non-protein helper molecules to catalyze their reactions. These are known as cofactors. Table 1 lists some important cofactors, and the dietary vitamins from which they are derived. Other important roles of vitamins, for example in activating molecules like Co-enzyme A and electron carrier molecules like NADH will be described later in the semester. A lack of a dietary source of a cofactor can often impair the activity of an enzyme.

#### *Integrated Control of Metabolism*

There are several ways in which enzymes are regulated, both based on intracellular and extracellular signals. An example might be that a lack of intracellular ATP causes an increase in ATP producing pathways such as glycolysis. On the other hand, low circulating blood glucose levels may work to stop a glucose consuming process such as glycolysis. We will discuss this in detail throughout the class, but some of the hormones we will discuss in this course that are particularly important are listed in Table 2:

Hopefully these hormones, how they work and how they are reg-

- <sup>11</sup> Some examples include:
- 1. Decreasing the activation energy
- 2. Increase the amount of the reactant(s)
- Decrease the amount of the product(s)
- 4. Increase the number of enzymes
- Decreasing the  $K_m$  of the enzyme
- 6. Shifting the substrate sensitivity of the allosteric enzyme

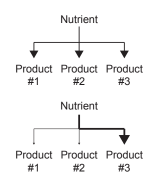


Figure 5: Example of how regulated pathways control nutrient fate.

Table 1: Some examples of cofactors that are important for enzymatic catalysis.

Cofactor	Source
TPP	Vitamin B <sub>1</sub>
Pyrdioxal Phosphate	Vitamin B <sub>6</sub>
Biotin	Vitamin H
THF	Folic Acid
Iron	Fe
Selenium	Se

Hormone	Main Function
Insulin	Reduces blood glucose and lipid levels
Adrenaline	Increases blood flow, nutrients to muscle
Glucagon	Increases blood glucose levels acutely
Cortisol	Increases blood glucose levels chronically
GH/IGF1	Promotes protein synthesis and bone growth
Testosterone	Promotes protein synthesis
Leptin	Suppresses appetite
CCK, Gastrin, Secretin	Regulation of digestion

Table 2: Some important metabolic hormones we will discuss in this class.

ulated is material you are familiar with from previous classes. If not, or you want a refresher, check out the Endocrine Control of **Macronutrient Metabolism** handout also available on Canvas.

## Post-Translational Modification

One common way by which enzymes are regulated is by the modification of existing proteins. One common example is protein phosphorylation. In this example a phosphate molecule is attached to an existing protein, which could increase or decrease its activity. This is often reversible, so a good analogy is that post-translational regulation is like flipping a switch for an enzyme on or off. This can occur fairly rapidly, and is not a permanent change.

#### Transcriptional Regulation

Another way to change the activity of a pathway is to selectively change the number of enzymes. If this is done at the messenger RNA level, it is known as transcriptional regulation. This is because transcription is the process by which new mRNA is made. By increasing or decreasing the rate of mRNA (and eventually protein) production, the cell can respond to a stimulus to make more or less of a particular protein. One example we will discuss in this course is that when cholesterol levels are low, the liver responds by making more cholesterol biosynthetic and retrieval enzymes and receptors. The regulation of transcription is often controlled by transcription factors, a class of proteins that can bind to selective sites of DNA and recruit the machinery to make (or prevent the making) of mRNA. These kinds of changes are slow, energetically costly and difficult to reverse. They represent a chronic response, and are not appropriate for short term modifications. The relationship between allosteric, post-translational and transcriptional regulation is demonstrated in Figure 6. Reflect on an example of metabolic regulation that you can think of. Then consider the timescale by which the changes happen, and try to think what would be the most appropriate mechanism to alter metabolism.

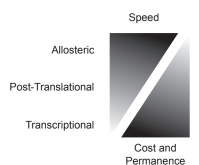


Figure 6: Shematic of the timing and permanence of some forms of enzymatic regulation.

# References

Bruce Alberts, Alexander Johnson, Julian Lewis, Martin Raff, Keith Roberts, and Peter Walter. Molecular Biology of the Cell. 4th edition, 2002. ISBN 0815332181.

J M Berg, J L Tymoczko, and L Stryer. Biochemistry, volume New York. 2013. ISBN 0-7167-3051-0.

Denise Ferrier. Lippincott Illustrated Reviews: Biochemistry. LWW, 1496344499, 7th edition, 2017. ISBN 1496344499.

Dara Leto and Alan R. Saltiel. Regulation of glucose transport by insulin: traffic control of GLUT4. Nature reviews. Molecular cell biology, 13(6):383-96, jun 2012. ISSN 1471-0080. DOI: 10.1038/nrm3351. URL http://www.ncbi.nlm.nih.gov/pubmed/22617471.