On Biology as an Emergent Science

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Abstract

Biology is considered here as an “emergent science” in the sense of Anderson and of Laughlin and Pines. It is demonstrated that a straightforward mathematical definition of “biological system” is useful in showing how biology differs in structure from the lower levels in Anderson’s “More is Different” hierarchy. Using cells in a chemostat as a paradigmatic exemplar of a biological system, it is found that a coherent collection of metabolic pathways through a single cell in the chemostat also satisfies the proposed definition of a biological system. This provides a theoretical and mathematical underpinning for Young’s fundamental model of biological organization and integration. Evidence for the therapeutic efficacy of Young’s method of analysis is provided by preliminary results of clinical trials of a specific application of Young’s model to the treatment of cancer cachexia.

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1 Why “Emergent Science”? 

1.1 Anderson, and Laughlin and Pines on “emergence”

In a famous paper Anderson[1] has pointed out that there is a natural hierarchy of scientific ideas. He starts with the usual (reductionist) strategy of the search for the laws obeyed by the elementary entities of physics, but then points out that the possibility of reduction does not imply constructivity. Rather, if science $Y$ underlies some science $X$: “The elementary entities of science $X$ obey the laws of science $Y$”. The $Y \rightarrow X$ hierarchy Anderson proposes is: elementary particle physics $\rightarrow$ solid state or many body physics; many body physics $\rightarrow$ chemistry; chemistry $\rightarrow$ molecular biology; molecular biology $\rightarrow$ cell biology; .... ; physiology $\rightarrow$ psychology; psychology $\rightarrow$ social sciences. Anderson goes on to state that “... this hierarchy does not imply that science $X$ is ‘just applied $Y$’. At each stage entirely new laws, concepts and generalizations are necessary, requiring inspiration and creativity to just as great a degree as in the previous one.” I heartily agree!

Although my conventional scientific career (in elementary particle physics) started out with the conventional scientific (reductionist) assumption that the only way to solve a basic scientific problem was to find the elementary entities, the laws they obey, and then construct higher levels of science from that basis, I now realize that I was mistaken. I have become convinced that 21st century science will be most exciting and fruitful if its basic problem is taken to be not only to find out if there are general hierarchy bridging laws that connect each level to the next and lead to novel types of complexity, but also if there are bridging laws which overarch the “elementary” bridging connection. This is one message I read into the paper on emergent science by Laughlin and Pines[2] who explicitly start from Anderson’s analysis. Laughlin’s book[3] is criticized by Leggett[4] under the title “Emergence Is in the Eye of the Beholder.” However, I am still particularly impressed by the fact that the values of $\hbar c/2e$ and $e^2/\hbar$ obtained by electric measurements in complex systems can be obtained to much higher accuracy than the values which can be obtained by direct “elementary particle” measurements, despite the fact that the details of the theories
used to understand the complex systems providing the data for these results have not reached consensus agreement.

1.2 The Organizing Principle of Darwinian Biology

Laughlin and Pines[2] also note that “For the biologist, evolution and emergence are part of daily life.” As Fred Young remarked when I started discussing emergent science with him, “Everything I ever said at ANPA (cf.[5]) was in the direction of emergent science”. This was sure to catch my attention because for several years Fred Young[6] has been trying to explain to me how his thesis work[7] is becoming more and more important for him as an explanatory tool of use in understanding how recent metabolic and physiological research all fits together. When James Lindesay joined these discussions of Fred’s work, our joint understanding began to take shape as a paper[8]. Briefly, I saw that Young’s results and Lindesay’s mathematical deduction from them could be interpreted as the starting point for adding the links “cell biology ↔ biological systems ↔ ecological systems and evolutionary biology” to Anderson’s proposed hierarchy in a specific way.

The careful reader will note that — in contrast to the earlier steps in Anderson’s hierarchy — I have used the symbol “↔” for the connective between levels of the hierarchy once biology enters the picture. The symbol “→” used by Anderson is an irreversible transition which replaces the “elementary particles” of the lower level by the laws they obey as the “elementary entities” of the new phenomena which occur at the more complex (higher) level, and which require the invention of new organizing principles, etc. which “emerge” from the careful study of this richer world of ideas. Biology arrived at its fundamental organizing principle by another route. Some biologists did not even believe that the phenomena they studied obeyed all of the laws of physics, in particular the second law of thermodynamics! Further, it was found useful to ask what purpose a particular aspect of these complex biological systems had “evolved” to satisfy. This kind of teleological explanation had been banished from physics after a very hard struggle, but its pragmatic usefulness in biology is hard to deny. That a “higher level” organizing principle can in fact lead
to *deductive* and *demonstrable* conclusions when applied “top down” to a lower level *biological* entity is one of the points I wish to make below. This is why I replace “→” by “↔” in the hierarchy once we enter the biological realm. Of course I must avoid the traps that lead to error when teleological reasoning is used carelessly. I hope the reader will reserve judgment as to whether I succeed in doing this until my methodology is presented clearly. In particular I believe that my methodology *also* avoids the traps pointed out by Anderson and by Laughlin and Pines, with whose basic conclusions I do agree.

Biology has sometimes been called a “Baconian” science in the sense that it started by amassing all kinds of details and facts assumed to be relevant to the subject and then induced general rules governing these facts. This methodology is to be contrasted with the tradition in the “mathematical” sciences which started from the astronomical practice of using numerical and geometrical models to make testable predictions. Skipping over vital historical details, this had the historical result that the “physical sciences” came to rely primarily on reductionism and hypothetical-deductive methodology for testing. Chemistry started out as a Baconian science, but began making the transition to a physical science in the nineteenth century thanks to electro-chemistry, thermodynamics and statistical mechanics. Quantum mechanics more or less allowed that transition to be completed; this transition has often been used as the leading example of the triumph of the reductive-hypothetical-deductive methodology.

Biology has not as yet made much *fundamental* use of mathematics. Its greatest nineteenth century success was the explanation of evolution via Malthus’ observation that (in a stable environment and over a sufficient period of time) a persistent population *must* have birthrate = deathrate. Starting from that deduction and with observations (in particular Darwin’s) of descent with modification, Darwin and independently Wallace came to the conclusion that “evolution by natural selection” is inevitable. This was the *non-quantitative* starting point for a scientific “evolutionary biology”. Since then this has remained unshaken as the organizing principle of *scientific* biology. Clearly — if my description of the history is roughly correct — this is a very different route to a basic organizing principle than the routes followed
in those “physical sciences” which now rest on hypothetical-deductive mathematical foundations.

One qualitative fact about biology which makes a methodological difference between it and the physical sciences is that “natural selection” inevitably presupposes the existence of some sort of environment within which the biological systems evolve, making it logically impossible to discuss biological systems without considering their interaction with that environment. I make explicit use of this fact in my definition of what is meant by a “biological system” in the sub-section which follows. A corollary of this point of view is that the paradigm of most importance in getting the study of biology off the ground is a persistent, evolved system. We will see that this provides a reference state, allowing fluctuations away from that reference state to be studied quantitatively.

1.3 A Proposed Mathematical Definition of a Biological system

I define a persistent biological system $B$ as a finite, countable population of individual constituents $C^B$ which in a suitable environment at constant temperature and pressure is a throughput (of molecules), steady state system satisfying the first and second laws of thermodynamics. The environment must supply food and fuel ($F$) at a rate sufficient to maintain the steady state. $B$ acts catalytically to convert the food and fuel into product molecules $P$ which are retained by the individual living constituents and waste molecules $W$ which are disposed of by the environment. The environment must also remove the waste heat required by the second law in such a way as to maintain the postulated constant temperature and pressure. The environment must remove that selection of living individual constituents whose disposal will maintain within the system (on average) a constant distribution of living constituents over all the states which can occur during the lifetime of any of them. The environment must absorb all dead constituents. This implies that “dead constituents” become part of the environment “at death” and are no longer part of $B$. In our context the (average)
number of (living) constituents satisfy the growth rate equation

\[ \dot{C}^B = k_B C^B \]  

and are said to be in a state of stable population (SP).

Note that \( C^B \) is a “counting number”. As such, it is necessarily dimensionless in terms of a physicist’s dimensional units of mass, length and time “\( M, L, T \)”. Then \( \dot{C}^B \) and \( k_B \) each have the dimension of inverse time (\( T^{-1} \)). By taking Eq. 1 as our defining equation for biology (in the context of a SP reference state), we emphasize in a different way the importance of the environment in our definition of biology. Lacking any evidence for a persistent physical environment, any biological system satisfying our definition — let alone its individual constituents — must have a finite lifetime.

2 The Young Model for the Organization and Integration of Biological Systems

2.1 The Chemostat as Paradigm for a Biological System

Our definition of product molecules \( P \) given in Sec. 1.3 allows us to specify the distribution of living constituents by the (average) number of product molecules they contain at any stage during the life cycle of each constituent. We illustrate how this can be done by narrowing the specific paradigm for a biological system used here to a group of cells in a chemostat maintained in a SP state. For the purposes of our theoretical analysis we assume that we can treat each cell in the chemostat as a coherent combination of its chemical constituents. Then we can use the symbol “\( C^B \)” to stand for a molecule in the chemist’s sense\(^\dagger\). This allows us to write chemical equations (conserving the numbers of each type of atom and the sum of their masses)

\(^{\dagger}\)A chemist’s “molecule” is a coherent structure which contains one or more chemical “atoms”, while a physicist usually thinks of an “atom” as composed of still more elementary constituents, and of a “molecule” as composed of two or more atoms.
connecting individual molecules to cells, which we take to be one of the (implicit) axioms of biochemistry.

We are now dealing with a population of growing cells inside the chemostat absorbing nutrient molecules $F$ and producing product molecules $P$ which are retained by the cell and waste molecules $W$ which are excreted into the solution surrounding the cell. Since the cell eventually divides into two cells which — at our level of analysis — are indistinguishable, we index the growing cells by the number of product molecules $n_P$ they have added in the range $N_P \leq n_P \leq 2N_P - 1$. Cell division is then the irreversible process

$$C^B(2N_P) \Rightarrow 2C^B(N_P)$$

(2)

The basic biochemical process in this context is

$$F + C^B(N_P + n_P) \Rightarrow C^B(N_P + n_P + 1) + W$$

(3)

Consequently the chemical equation describing the operation of the chemostat in this simplest case is

$$N_F F + \sum_{n_P=0}^{2N_P-1} C^B(N_P + n_P) \Rightarrow \sum_{n_P=1}^{2N_P-1} C^B(N_P + n_P) + 2C^B(N_P) + N_W W$$

(4)

which, by defining a complete population of cells (i.e. a population which, in the appropriate environmental context, when supplied with $N_F$ nutrient molecules, can produce two clones by cell division) as $\Sigma C^B \equiv \sum_{n_P=0}^{2N_P-1} C^B(N_P + n_P)$, we can rewrite as

$$N_F F + \Sigma C^B \Rightarrow \Sigma C^B + C^B(N_P) + N_W W$$

(5)

or as

$$N_F F \Rightarrow \Sigma C^B C^B(N_P) + N_W W$$

(6)

A growing cell has to add $N_P$ product molecules to its structure before it can divide and start the process over again. One of those two copies (clones) must be removed
at some subsequent time (in its life cycle or when it dies); this pruning is required to maintain the SP state. In this particulate description of the overall process, any growing cell will have to add each individual product molecule sequentially. We assume that the context in which the equations apply is a through-put steady state (SP state). Then the rate at which the molecules of $F$ move into the growing cell, the rate at which the molecules of $P$ join the growing cell, the rate at which the cell divides into two clones (beginning cells), and the rate at which one of these two growing cells is eventually pruned from the cell colony are all the same. That is

$$[\dot{C}_B] = k_B[C^B]; \quad [\dot{F}] = k_B[F]; \quad [\dot{P}] = k_B[P]; \quad [\dot{W}] = k_B[W] \quad (7)$$

Here the symbol $[X]$ ($X \in C^B, F, P, W$) means the concentration (i.e. mass per unit volume) of the substance $X$. For small molecules (i.e. molecules whose atomic content and (if needed) molecular structure are known) this mass is most conveniently measured in terms of moles (i.e. gram molecular weights). These equations immediately suggest that it may be possible to treat concentrations of small molecules as biological systems in an appropriate context. We develop this idea in the next sub-section, in which we give precision to the concept of metabolic pathway.

The careful reader will have noted that we have use the symbol $\Rightarrow$ denoting the irreversibility of the chemical reaction not only for cell-division (Eq. 2) but also for the individual step (Eq. 3) in which the cellular environment catalyzes the transition from food molecule(s) to the product and waste molecules. We assume that this can only happen when the initial and final molecules are in the correct stoichiometric ratios (see next section). This is because we are interested in this paper only in the passage of molecules through the cell (or to their location within the cell) when this path does go through some catalytic site (which we will call an enzyme) that guarantees that we are talking about a throughput steady state which is far from equilibrium and not about the equilibrium states with which much of physical chemistry is concerned. Thus there are no two-way transitions at the basic level and the usual use of detailed balance rate constants is, from the start, inapplicable. This brings us to the discussion of (enzymatic) metabolic pathways in the next section.
2.2 A Coherent Collection of Metabolic Pathways as a Paradigm for a Biological System

The food/fuel molecule or molecules $F$ that initiate the basic process (Eq. 3) could have entered the cell at many different places, and the waste molecule or molecules that complete the process can leave the cell at many different places, but (in our abstract model) the critical transition occurs at only one place along the path(s) connecting the input and output surface patches, namely where some enzyme $E_{F \rightarrow PW}$ catalyzes the reaction $h_F F \Rightarrow h_P P + h_W W$. We call this “one dimensional” route through the cell a metabolic pathway and represent its action by the biochemical equation

$$E \cdot h_F F$$

$$\uparrow \quad \downarrow$$

$$h_F F + E \quad E + h_P P + h_W W$$

$$\downarrow \quad \uparrow$$

$$E$$

Eq. 8 represents the irreversible, catalytic action of a single enzyme molecule, which may dynamically change its shape during the process but automatically resumes its initial shape after the process is completed\(^1\). Note that, for the biochemical processes used in our paradigm, this process must occupy a (3+1)-dimensional space-time volume and hence must be nonlocal. The numbers $h_F, h_P$ and $h_W$ must be integers because both the number of (chemical) atoms and the amount of (chemical) mass are conserved in the process. Their ratios $h_{X/Y} \equiv h_X / h_Y = (h_Y / h_X)^{-1} = h_{Y/X}^{-1}; X, Y \in F, P, W, \ldots$ are called stoichiometric ratios. If we measure the concentration $[X]$ [which has physical dimensions $ML^{-3}$ (i.e. mass per unit volume)] of any chemical substance $X$ in moles (i.e. in gram-molecular weights per unit volume), then

\(^1\)This restoration of the initial state of the enzyme provides one “feedback” control mechanism. Some feedback control loop in the information flow is required for any persistent, self-organizing complex system to exist.
the stoichiometric ratios are identical to the concentration ratios. Then the equation also can be read as the number of moles of each substance which will react in this way when catalyzed by one mole of the enzyme. Note that we can now rigorously and quantitatively bridge the small molecule $\leftrightarrow$ cell mass magnitude gap by writing, as a corollary to Eq. 6

$$h_F N_F = h_P N_P + h_W N_W$$

(9)

Note that this is an algebraic equation connecting positive definite integers and is not a chemical equation.

A few comments are needed here. Note that the $N_P$ apparently independent metabolic pathways implied by Eq. 4 — which are needed in order to allow Eq. 7 to be treated as defining the hierarchical nesting of a collection of biological systems — must act coherently, at least at the conceptual level; this assumption is also needed in order for the cell to be thought of as a coherent chemical molecule. The conceptual advantage of this step is to allow the very complicated process of cell growth and division to be made into the simple doubling of the starting cell via the sequence of steps (Eq. 4) that leads to cell division (Eq. 2). Then the rate $k_B$ at which the transition occurs is a quantitative and experimentally measurable function of the concentrations of small molecules of known structure called here $F$, $P$ and $W$, even if we do not know the molecular weight of the enzyme invoked by Eq. 8, or the details of how the catalytic result is achieved, let alone knowing the molecular weight of the cell!

Some such critical conceptual step is needed in order for the mathematical model we are constructing to be able to explain how chemostats can determine empirically what function of these concentrations the cell growth rate $k_B$ is. That such functions are known is an empirical fact[9]. It is this fact which allows us to go from it to a simple mathematical formulation of Young’s model. Explicitly we quote from Fred’s thesis ([7], p.1)

\[ \ldots \text{the value of } k_B \text{ is a reproducible function of the medium composition}[9] \]

\[ \ldots \]
which we write formally as

\[ k_B = K_B([F_1], [F_2], ..., [F_j], ..., [F_J]) \] (10)

Here the nutrients \( F_j \) are distinguished from each other by the unique enzymes \( E_j \) which catalyze the irreversible reactions

\[ hF_jF_j \xrightarrow{E_j} hP_jP_j + hW_jW_j \] (11)

that remove \( hF_j \) molecules of \( F_j \) from the metabolic pathway and replaces them with product \( (P_j) \) and waste \( (W_j) \) molecules, conserving chemical mass and atom flux. \( J \) is the number of types of enzymes and the number of types of metabolic pathways we consider important in any particular analysis. \( K_B \) is not a function in the usual mathematical sense. For us, if the values of the parameters are known over the ranges of values and to the accuracy needed for our immediate purposes, a “table lookup” plus any well defined “interpolation procedure” suffice to make this framework into a testable theory in Popper’s sense.

Accepting that Eq. 10 is a reproducible empirical statement based on table lookup has important consequences. In that context the inescapable fact is that all the numerical quantities (in this case \( k_B \) and each of the \([F_j]\)) have an experimental range of uncertainty. I formalize this fact by assuming that whenever we assert Eq. 10, we are claiming that there are \( 2(J + 1) \) numbers called \( k_{\text{min}}, k_{\text{max}}, [F_j]_{\text{min}}, [F_j]_{\text{max}} \) such that for any choice of numerical values within these ranges, no matter how correlated, the asserted equality provides an acceptable representation of the data for our purposes. If this statement becomes suspicious, the careful experimenter will look for an explanation either in some source of systematic error, or some theoretical constraint or possibility that has been ignored. Note that in either case, these limits become testable hypotheses in Popper’s sense, and new experiments can either reduce the experimental uncertainty or produce new empirical knowledge. This is standard procedure in physics.

With this understood, we can use some hypothesis that makes nutrient “\( j \)” the “most important” for the purposes of our analysis, and formally “invert” Eq.10 by
defining

\[ F_j = (K_B)^{-1}_j(k_B; [F_1], [F_2], ..., [F_{j-1}], [F_{j+1}], ..., [F_J]) \approx (K_B)^{-1}_j(k_B) \]  \hspace{1cm} (12)

which means that, to the extent that the approximation is valid, we can ignore what is going on in the concentrations of the other nutrients and find some monotonically increasing function of \( k_B, k_{min} < k_B < k_{max} \) to fit the observed values of the correlated variation of \( [F_j] \), for \( [F_j]_{min} < [F_j] < [F_j]_{max} \) (or \( visa versa \)), — i.e. \( k_B = (K_B)_j([F_j]) \).

With this basic phenomenology understood we can make testable empirical hypotheses about and place reasonable theoretical constraints on the concept of “metabolic pathway” in the context of a stable population of bacterial cells in a chemostat.

2.3 Single Enzyme Control in a single metabolic pathway as an irreversible transition

The simplification of Eq. 8 for each enzyme/pathway \( j \) to Eq. 11 allows us to compare it to the detailed model for catalytic action in the irreversible reaction

\[ 2 \text{CO} + O_2 \xrightarrow{\text{Cat.}} 2 \text{CO}_2 \]

as analyzed by Grinstein, et. al.[10]. As the authors note,

Since the reverse reaction \( \text{CO}_2 \rightarrow \text{CO} + O \) is not allowed, the system defined by the above rules cannot satisfy detailed balance for any underlying Hamiltonian.

which reinforces the remark already made at the end of sub-section 2.1 that the processes we are considering cannot be described by the rules of equilibrium physical chemistry. It also warns us (in our non-equilibrium context of irreversible, steady state, throughput processes) that we cannot expect the essential mathematics needed for theoretical biology to resemble the continuum mathematics used in classical theoretical physics. I fear this fact about biological systems is often ignored by biochemists analyzing enzyme reactions \( in \text{ vitro} \). The advantage Young has in basing his model
on chemostat data is that these empirical studies are, in fact, \textit{in vivo} experiments. They allow us to go \textit{directly} from chemical measurements (concentrations of small molecules) to a parameter ($k_B$) that measures the (average) time it takes a living organism to replicate itself in an environmental context that allows a biological system composed of such organisms to achieve a persistent steady state (SP-state).

The rules the authors\cite{10} refer to in the quote given above describe the way to parameterize the rates at which the incoming molecules attach to the catalytic surface, rearrange bonds to form the product molecules of the outgoing gas, and the rates at which the outgoing molecules detach. These details need not concern us here, nor do the numerical methods Grinstein, et. al. are forced to use because they lack a Hamiltonian model. What does concern us is that the catalyzed transition \(2CO + O_2 \Rightarrow 2CO_2\) is a worked out example analogous to the way the $F$ molecules come along the \textit{incoming} part of the metabolic pathway to a specific enzyme and the $P$ and $W$ molecules leave on the distinctly different \textit{outgoing} part of the same pathway. We \textit{could} make a more detailed model of this process, but we are \textit{not required} to do so in order to achieve our results. All we need abstract from the complicated process that goes on in the ill-defined space-time volume around the enzyme is the fact that this transition separates the metabolic pathway into an incoming and an outgoing part, and that it \textit{fixes} the stoichiometric ratios of all the substances in this single metabolic pathway whose mass flow is continuous through this volume.

The reason we are not concerned with the geometrical details is that the basic equations (Eq. 7) are \textit{space-scale invariant} and only depend on spatial averages (concentrations) as functions of time. To smooth these out in the complicated region where $F$ attaches to the enzyme, the enzyme rearranges $F$ into $P$ and $W$ and these leave, we assume this region has an average length $L$. We assume that an average flow velocity for molecules along this metabolic pathway through the cell can be defined by $v = k_B L$. Then within this region, we can measure distance along the pathway by a spatial coordinate $x = vt = Lk_B t$ when $0 < x < L$, $0 < t < T_B = k_B^{-1}$. Assume that the pathway is in an SP-state (i.e. $v = \text{const.} = k_B L = L/T_B$). \textit{Upstream} of this transition region (i.e $x < 0$) the concentration $[F_j]$ must have a constant \textit{input} value which we call $[F_j]_I$. \textit{Downstream} of the transition region the concentrations $[P_j]$ and
[W_j] must have constant output values which we call [P_j]_O and [W_j]_O. We then know that the concentration [F_j] must fall from its input value [F_j]_I to zero as it passes through the transition region. This shows that we can always describe the steady state action of any enzyme which causes an irreversible phase transition by

\[ \dot{F}_j = -k_B[F_j] \]  

(14)

if we use the algebraic sign conventions a) that \( k_B \) is a positive definite constant and b) that the time rate of change of a concentration is positive when it is the same as the sign of the rate change of a growing cell. Similarly [P_j] must start at zero at \( x = 0 \) and rise to the output value [P_j]_O at \( x = L \). Then a little thought tells us that if Eq. 8 is used to represent an irreversible transition, we must have that

\[ \dot{Y}_j = k_B[Y_j] \]  

(15)

Now we must face up to the fact that, empirically, the chemostat data often exhibit a substantial range of values of \( k_B \), as is implied by Eq. 10 and the quotation which it formalizes. Indeed, the question of how such a strictly correlated (by any set of values or range of values for \( k_B \) for which the quote and or Eq. 10 are a correct representation of the facts) can come about was the problem Fred Young’s thesis set out to solve.

Here the Darwinian organizing principle of natural selection comes to our aid. Any organism in a biological system will benefit by extending the range of the concentrations of nutrients which it can tolerate and continue to reproduce, and the speed with which it can make use of them in competition with other organisms or with genetically modified members of its own species. But the metabolic pathways within each organism (having the same genome) will gain collectively (for its genotype) if their action is tuned to make maximum use of the total supply which can be absorbed by the organism as a whole. On both counts we expect an organism-wide coordination to be selected for, and not just maximum range and efficiency of the action of the individual catalytic pathways. As is not surprising, from the point of view of the central dogma of molecular biology, this coordination is provided by the
genetic control of the production of the enzymes themselves. Since this is more easily explained by using the control mechanism discovered by Fred Young in his thesis than by discussing a single enzyme pathway, we now turn to that explanation in the next sub-section.

2.4 Two Enzymes linked by a feedback loop in a single metabolic pathway

The basic feedback control loop for metabolic regulation which Fred Young[7] discovered, written as a chemical equation, is

\[
\begin{array}{c}
h_UU \\
\uparrow \quad \downarrow \\
\downarrow \quad \uparrow \\
h_ID \\
\end{array}
\]

\[
\begin{array}{c}
h_I I + E_I \\
\downarrow \\
h_D D \\
\end{array} \quad \quad \quad \quad \quad \begin{array}{c}
E_O + H_O O \\
\uparrow \\
\downarrow \\
h_UU \\
\end{array}
\]

Note that this connects two irreversible enzyme-catalyzed transformations, namely

\[
h_I I + h_DD \Rightarrow E_I \quad h_UU \Rightarrow h_DD + h_OO
\]

Note that D — which is analogous to the the enzyme E in Eq. 8 (if we replace E \cdot F by D \cdot I = U) — is conserved in the sense that it retains (in a SP-system) a constant concentration. Note also that (in a SP-system) the net effect is to transform I to O irreversibly, conserving mass and chemical atoms, at a rate

\[
k_B = \frac{\dot{O}}{[O]} = -\frac{\dot{I}}{[I]} = \frac{\dot{U}}{[U]} = \frac{\dot{D}}{[D]}
\]

This is, of course, the same conclusion we reached about the action of the individual enzymes (cf. Eq.’s 14 nd 15). This means that we can go on connecting nodes (representing enzymes which unequivocally direct mass flow in the direction defined by positive growth rate or, in feedback links, unequivocally in the opposite direction)
in a way that will never upset the SP character of the system provided the environment is stable and we can prove that the system is stable against “normal” fluctuations in the environment.

Further, Eq. 18 implies that \[ \frac{\dot{O}}{\dot{I}} = -h_{O/I}, \] the negative of the stoichiometric ratio of the concentrations. Not only is the rate of decrease of I precisely equal to the rate of increase of O (a fact we could derive directly from chemical mass and atom conservation), but if we think of the cell as a factory for the production of O, ratios of the stoichiometric coefficients in Eq. 16 could serve as the set-points for some rate control system that optimizes the use of resources I to the rate at which they are provided. This is obvious to a chemical engineer. That natural selection has “engineered” such a system is a deduction from the Darwinian organizing principle.

Fred Young’s approach is to “reverse-engineer” the data on the concentrations in steady state throughput experiments using what is known about the structure and working of the cell so as to tease out how the control system operates. One advantage of using his control cycle, rather than concentrating on the genes (and hence the enzymes) directly, is that his control loop allows this to be done using only the concentrations of the small molecules as the empirical starting point. As he notes ([7], p.8): “The interrelationships between cellular components that define the steady-state and illustrate the scope of regulation which is independent of specific [genetic] induction-repression mechanisms have been comprehensively tabulated.”

One general mechanism for cell-wide rate control recognized by Young is the relation between protein synthesis and ribosome synthesis when both are thought of as a function of \( k_B \) (cf. [7], Fig. 7, p.37 and related text). For a stable population of growing cells, and a large range of values of \( k_B \), the rate of protein synthesis (per genome equivalent of DNA) is constant, whereas the relative rate of ribosome synthesis is a rapidly increasing function of \( k_B \). Clearly the value of \( k_B \) where these two curves cross is a “rate control set point”.

Why is this true? The proteins are manufactured by the ribosomes. The particular protein called for is coded on an “instruction tape” (messenger ribonucleic acid — mRNA). This tells the ribosome which of the 20 possible amino acids to attach next onto the growing protein (polypeptide) chain. An “expressed” DNA-gene uses
one of 20 “three letter codons” (corresponding to the 20 amino acids which can be used to make a protein chain) to provide the information added sequentially to the mRNA instruction tape. The fact that the concentrations of ribosomes, ribosomal nucleic acid (rRNA), mRNA and tRNA are all proportional to $k_B$ then tells us (accepting the one gene - one enzyme doctrine and still subtler approximations) that we can expect the concentration of any enzyme (a “large molecule” made up of one or more polypeptide (protein) chains) produced by this machinery to also be proportional to $k_B$. The fact that the (relative) rate of protein synthesis as a function of $k_B$ is constant (in the range where it crosses the rate of ribosome production) can be interpreted as due to the likelihood that the rate of transcription for any of the codons that specifies any amino acid is approximately the same as for any other codon.

The next step is to note that the amount of the enzyme synthesized is controlled by the expression of the gene and that this, in turn, is controlled by the operator-promoter region of the gene. These controls can be either positive (enzyme induction) or negative (enzyme repression) and can be effected by a change in the concentration of any appropriate small molecule or protein in the metabolic pathway upstream of the enzyme in question, even though it is not directly involved in the $I \Rightarrow O$ catalysis. [Such a change can even “turn off” the gene completely and hence form the starting point for a concentration threshold-controlled on-off “switch”. We will mention such switches in the discussion of the cancer cachexia treatment, but not model them in this paper.]

\[\textit{DNA is a shorthand for deoxyribonucleic acid, The transfer RNA (tRNA) — with 20 varieties — transfers uniquely the sequential information from an expressed mRNA transcript of a DNA “structural” gene one codon at a time by having one end which attaches by complementary base pairing to the RNA codon and picks up on the other end the cognate amino acid which is added to the growing polypeptide chain. If this apparatus worked perfectly there would be approximately one “constant” rate (with a fine structure of 2 or 20 or some number less than 64 rates) for this process. But the mRNA can itself get degraded at some rate between the time when the information is transferred to it physically and the time when it is read. Consequently information transfer and the transfer of the material coding of that information can have different average rates. Fortunately, for the purposes of this paper, we can ignore these complexities.}\]
Two applications of this control loop are discussed in [7]. The first is glucose metabolism. For it “I” is some phosphate in the food which is picked up by adenosine-diphosphate (ADP) [identified with “D”] to form adenosine-triphosphate (ATP) [identified with “U”], which then passes on the phosphate to some downstream product [identified with “O”] and returns ADP as the feedback which completes the cycle. Thus there is an internal “set-point” for the internal control loop — namely the stoichiometric ratio $h_{U/D}$ — and an external set-point, $h_{I/O}$. Both are under the genetic control of the two genes associated with the two enzymes $E_I$ and $E_O$. The second application is the addition of a monomer as the next link in the chain of a growing polymer. The second example is illustrated in [7], Fig. 5, p.28, which is reproduced below, together with its figure caption as Figure 1. I trust that, after the above sketch of how the whole thing works, this captioned figure is self-explanatory.

What Fig. 1 does not point out is that the genetic control mechanism acts on a long time scale, appropriate to a secular change in the rate and/or amount at which food is flowing into the system from the environment, while the internal control loop acts on a shorter time scale which can smooth out short term fluctuations in the concentrations due to other causes. That this feedback is stable follows immediately from the irreversibility of the direction of flow at the two enzymes. These absorbing state phase transitions (from $M + X$ to $(X \cdot M)$ and from $(X \cdot M)$ to $X + P$ — with the return control loop path that takes $X$ “back” to the initiating phase transition in place) also enforce the stoichiometric set points for both the interior and the exterior rate control. The rate controlled range of $k_B$ comes from: a) the fact that the number of ribosomes per cell increases with $k_B$; b) the fact that the range of $k_B$ is limited at the upper end by the maximum number of ribosomes which the cell can hold in a steady state; c) the fact that the range of $k_B$ is limited at the lower end by the minimum amount of nutrient which will allow, at least, the minimum stable number of cells to maintain themselves in the chemostat at the flow rates for nutrient input and for the solvent carrier input.
Figure 1: Representation of how monomer charge can be maintained at various steady-state growth rates through the use of rate effectors. When rates of monomer producing and utilizing reactions are balanced the concentration of \((\text{X} \cdot \text{M})\) is proportional to the rate of synthesis of \((\text{X} \cdot \text{M})\). \((\text{X} \cdot \text{M})\) is a positive effector for synthesis of both \(\text{X}\) and \(\text{P}\). In this way the rates of synthesis of \(\text{P}\) and \(\text{X}\) will be proportional to the rate of synthesis of \((\text{X} \cdot \text{M})\). During non-steady-state conditions the concentration of \((\text{X} \cdot \text{M})\) will no longer be proportional to the rate of synthesis of \((\text{X} \cdot \text{M})\) and consequently rates of synthesis of \(\text{P}\) and \(\text{X}\) will no longer be equal to the rate of monomer availability. \(\text{P/O}\) is the operator promoter region of the structural gene. \(\text{X}\) represents a carrier of monomer units. (The thickness of the arrow is not meant to reflect the relative reaction rates.)

The general applicability of Fred Young’s general in-out, up-down rate control loop feedback model, i.e.

\[
\begin{array}{c}
U \\
\uparrow \downarrow \\
\rightarrow I \quad O \rightarrow \\
\downarrow \uparrow \\
D
\end{array}
\]  

(19)

to most (possibly all) biological systems, and to many well-modeled physical systems that provide significant analogies for biological systems, may not be obvious. After all it was only first discovered in \textit{E-coli} metabolism. But careful perusal of Young’s
thesis[7], should begin to remove doubt on that score. The thesis was deliberately undertaken, not to solve a specific problem but to find a mechanism that could account in a general way for how rate control of metabolism can lead to multiple rates of growth in stable populations and for growing systems exhibiting balanced exponential growth. Subsequent developments in biology and other fields provide ample evidence for the fact that Young’s model is ubiquitous in its applicability. This topic will be discussed in[11], where the connected chain: non-equilibrium steady state → absorbing state phase transition → allometric scaling laws → fractal scaling → Kolmogorov scaling is developed. The abstract of an earlier talk by Young on this subject at an international meeting in Shanghai is presented here as Appendix 1.

Although the primary purpose of this paper was to prepare a mathematical and logical basis for the Young model, whose technical structure will be presented elsewhere[8], we wish to also take this occasion to point out that, once the biological principles are understood, the top-down analysis of metabolic pathways which Young’s general model makes possible did not have to wait for mathematical development in order to be applied. The underlying logic and analytic framework have been used as the basis for research to identify, select and assemble data from the published literature. This data can then be used to create models for Cancer Cachexia and other diseases. This systematic approach, starting from Young’s 1977 thesis[7], developed by Young and collaborators and now called HiNET, then allows combined drug therapies appropriate to these diseases to be constructed. The technology has led to a venture capital backed company with a portfolio of high potential ideas, one already in FDA-approved Phase 2 trial and two more likely to enter trial in 2008.

The problem of cancer cachexia can be briefly described as follows. Normal nutrition for our species and many others has a replete-hungry cycle with on-off switches changing the metabolic pathways between the two stable states. In certain shock states, there a great need for nutrition at any cost and the body in these states starts eating anything inside it, including its own structure. Normally this state turns off when the danger is past, but cancer and some cancer therapies can produce a shock state that does not return to normal; consequently the body wastes away even though ample nutrition is provided by injection into the veins. Using his analysis,
Young found a way to treat the patient with combinations of FDA-approved drugs. They alter the concentrations of small molecules in the direction which returns the body to normal nutritional states and solves the problem. Preliminary clinical trials were successful and second stage trials were approved. An older short report of this is given in Appendix 2.

In conclusion, I believe that Young’s control loop feedback in-out: up-down cycle model for a throughput system is a good candidate to become an emergent fundamental law of biological systems going beyond the Darwinian organizing principle. Using such control loops as coupled nodes in a hierarchical model for top-down analysis of functional metabolic pathways, of which the first examples are Young’s HiNET models, bids fair to become a fruitful research tool for uncovering novel emergent biological organizing principles during the 21st century.

3 Acknowledgments

This paper rests primarily on the decades of work by Fredric S. Young on his unique approach to the problem of the organization and integration of biological systems. I am most indebted to him for his invitation to participate in this research. I am also indebted to James V. Lindesay for his collaboration on clarifying the mathematical structure of the feedback system involved in the two-enzyme control loop, and to both of them for the three-way discussions we had during 2004-2006. I owe much to the clarification of the logical and philosophical structure of what we are attempting to do provided by Walter R. Lamb while he was still with us.

4 Appendix 1: The Universal Modular Organization of Hierarchical Control Networks in Biology

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Progress in the physical sciences have always involved conceptual and theoretical simplification and unification. Modern biology has resisted this tendency and has focused almost completely on the details. The sequencing of the human genome has not been translated into comprehensive models and has not led to new therapies. Using reverse engineering, we have abstracted a theoretical description of the universal modular organization of biological control systems which are modeled as the construction of a fractal representing the hierarchical control network or HiNet. Disease therapy becomes a problem of shifting the state of the HiNet to a configuration closer to normal homeostasis. This has enabled the rational and systematic development of combination therapies for clinical trials. An emphasis on energy and control manifolds connects this approach to catastrophe theory. The modularity of HiNet allows a hierarchical network decomposition and modeling of local processes on low dimensional control manifolds. Modeling of the integrated global organization of a biological system requires control spaces of many more dimensions than 3 as stated by Thom in *Structural Stability and Morphogenesis*. A HiNet model of allometric scaling supports the recent application by Ji-Huan He of El Naschie's $\varepsilon^\infty$ theory to biology. (Abstract of paper presented at the 2005 International Symposium on Non-Linear Dynamics: Celebration of M.S. El-Naschie’s 60 Anniversary, December 20-21, Shanghai, China)

5 Appendix 2: The Obsolescence of Reductionist Biology: Systems Biology Modeling and Cancer Cachexia Therapy Development Based on Emergent Patterns of Organization Rather Than on Genes and Molecules

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Vicus has developed a hierarchical network (HiNET) model of emergent patterns of organization based on principles of self-organized criticality, phase-transitions, in-
tegral control and reaction blocks. We will describe our HiNET model of cancer cachexia, a catastrophic wasting disorder secondary to advanced cancer, and its predicted EKG-based biomarkers and reaction-block drug targets. We will show data from our retrospective and prospective VT-122 clinical trials and contrast our clinical results with previous failed attempts targeting specific dysregulated pathways and proteins. (Abstract of paper presented at Beyond the Genome 2006: Top Ten Opportunities in the Post-Genome Era, June 19=21, 2006, Fairmont Hotel, San Francisco, California.)

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23
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