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Preface

The aim of the *Advances of Biochemical Engineering/Biotechnology* is to keep the reader informed on the recent progress in the industrial application of biology. Genetical engineering, metabolism and bioprocess development including analytics, automation and new software are the dominant fields of interest. Thereby progress made in microbiology, plant and animal cell culture has been reviewed for the last decade or so.

The Special Issue on the History of Biotechnology (splitted into Vol. 69 and 70) is an exception to the otherwise forward oriented editorial policy. It covers a time span of approximately fifty years and describes the changes from a time with rather characteristic features of empirical strategies to highly developed and specialized enterprises. Success of the present biotechnology still depends on substantial investment in R & D undertaken by private and public investors, researchers, and entrepreneurs. Also a number of new scientific and business oriented organisations aim at the promotion of science and technology and the transfer to active enterprises, capital raising, improvement of education and fostering international relationships. Most of these activities related to modern biotechnology did not exist immediately after the war. Scientists worked in small groups and an established science policy didn't exist.

This situation explains the long period of time from the detection of the antibiotic effect by Alexander Fleming in 1928 to the rat and mouse testing by Brian Chain and Howart Florey (1940). The following developments up to the production level were a real breakthrough not only biologically (penicillin was the first antibiotic) but also technically (first scaled-up microbial mass culture under sterile conditions). The antibiotic industry provided the processing strategies for strain improvement (selection of mutants) and the search for new strains (screening) as well as the technologies for the aseptic mass culture and downstream processing. The process can therefore be considered as one of the major developments of that time what gradually evolved into "Biotechnology" in the late 1960s. Reasons for the new name were the potential application of a "new" (molecular) biology with its "new" (molecular) genetics, the invention of electronic computing and information science. A fascinating time for all who were interested in modern Biotechnology.

True gene technology succeeded after the first gene transfer into *Escherichia coli* in 1973. About one decade of hard work and massive investments were necessary for reaching the market place with the first recombinant product. Since then gene transfer in microbes, animal and plant cells has become a well-

established biological technology. The number of registered drugs for example may exceed some fifty by the year 2000.

During the last 25 years, several fundamental methods have been developed. Gene transfer in higher plants or vertebrates and sequencing of genes and entire genomes and even cloning of animals has become possible.

Some 15 microbes, including bakers yeast have been genetically identified. Even very large genomes with billions of sequences such as the human genome are being investigated. Thereby new methods of highest efficiency for sequencing, data processing, gene identification and interaction are available representing the basis of genomics – together with proteomics a new field of biotechnology.

However, the fast developments of genomics in particular did not have just positive effects in society. Anger and fear began. A dwindling acceptance of “Biotechnology” in medicine, agriculture, food and pharma production has become a political matter. New legislation has asked for restrictions in genome modifications of vertebrates, higher plants, production of genetically modified food, patenting of transgenic animals or sequenced parts of genomes. Also research has become hampered by strict rules on selection of programs, organisms, methods, technologies and on biosafety indoors and outdoors.

As a consequence process development and production processes are of a high standard which is maintained by extended computer applications for process control and production management. GMP procedures are now standard and prerequisites for the registration of pharmaceuticals. Biotechnology is a safe technology with a sound biological basis, a high-tech standard, and steadily improving efficiency. The ethical and social problems arising in agriculture and medicine are still controversial.

The authors of the Special Issue are scientists from the early days who are familiar with the fascinating history of modern biotechnology. They have successfully contributed to the development of their particular area of specialization and have laid down the sound basis of a fast expanding knowledge. They were confronted with the new constellation of combining biology with engineering. These fields emerged from different backgrounds and had to adapt to new methods and styles of collaboration.

The historical aspects of the fundamental problems of biology and engineering depict a fascinating story of stimulation, going astray, success, delay and satisfaction.

I would like to acknowledge the proposal of the managing editor and the publisher for planning this kind of publication. It is his hope that the material presented may stimulate the new generations of scientists into continuing the rewarding promises of biotechnology after the beginning of the new millenium.

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The Morphology of Filamentous Fungi

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The morphology of fungi has received attention from both pure and applied scientists. The subject is complicated, because many genes and physiological mechanisms are involved in the development of a particular morphological type: its morphogenesis. The contribution from pure physiologists is growing steadily as more and more details of the transport processes and the kinetics involved in the morphogenesis become known. A short survey of these results is presented.

Various mathematical models have been developed for the morphogenesis as such, but also for the direct relation between morphology and productivity – as production takes place only in a specific morphological type. The physiological basis for a number of these models varies from thorough to rather questionable. In some models, assumptions have been made that are in conflict with existing physiological know-how. Whether or not this is a problem depends on the purpose of the model and on its use for extrapolation. Parameter evaluation is another aspect that comes into play here.

The genetics behind morphogenesis is not yet very well developed, but needs to be given full attention because present models and practices are based almost entirely on the influence of environmental factors on morphology. This makes morphogenesis rather difficult to control, because environmental factors vary considerably during production as well as on scale. Genetically controlled morphogenesis might solve this problem.

Apart from a direct relation between morphology and productivity, there is an indirect relation between them, via the influence of morphology on transport phenomena in the bioreactor. The best way to study this relation is with viscosity as a separate contributing factor.

Keywords. Environmental factors, Filamentous fungi, Genetics, Modelling, Morphology, Physiology, Transport phenomena

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List of Symbols and Abbreviations

C	Concentration, kg m^{-3}
C_x	Concentration of biomass, kg m^{-3}
DCR	Diffusion with chemical reaction
ID	Diffusion coefficient, $\text{m}^2 \text{s}^{-1}$
DOT	Dissolved oxygen tension, N m^{-2}
D_r	Stirrer diameter, m
d_h	Diameter of hypha, m
ER	Endoplasmatic reticulum (an internal structure element of a cell)
$f(x, t)$	Population density function: number per m^3 with property x at time t
k_1, k_2	Lumped parameters
k_{1a}	Mass transfer parameter, s^{-1}
L	Length of hypha, m
L_e	Length of main hypha in hyphal element, m
$L_{e\text{max}}$	Maximum length of main hypha capable of withstanding fragmentation, m
L_{equil}	Equilibrium length, m
L_t	Length of all hyphae in hyphal element, m
L_{hgu}	Length of hyphal growth unit (L_t/n), m
m	mass, kg
m_{hgu}	Mass of a hyphal growth unit, kg per tip
N	Rotational speed of stirrer, s^{-1}
n	Number of tips in hyphal element, -

NADP	Nicotinamide adenine dinucleotide phosphate: oxydation/reduction coenzyme in which NADPH is the reducing substance
P/V	Power per unit volume of fermenter, $W m^{-3}$
r	Distance to stirrer, m
r(C)	Reaction rate as function of C, $kg m^{-3} s^{-1}$
r_l	Rate of vesicle production per unit length of hypha, number $m^{-1} s^{-1}$
Rho 1p	A GTP-binding enzyme involved in the cell awl synthesis
t_c	Circulation time, s
V	Volume, m^3
v	Velocity, $m s^{-1}$
V_{disp}	Volume with maximum dispersion potential, m^3
z	Vector representing the environmental conditions, varying dimensions
ϵ	Power per unit mass, $W kg^{-1}$
ϕ_p	Pumping capacity of stirrer, $m^3 s^{-1}$
γ	Shear rate, s^{-1}
μ	Specific growth rate, s^{-1}
τ	Shear stress, $N m^{-2}$

1

General Introduction

Filamentous fungi are fascinating organisms, not only because of the inherent beauty of their fruiting bodies but also because of their complicated and scientifically very interesting behaviour. They are also able to produce a large variety of useful, commercially interesting products.

The use of filamentous fungi as production organisms in industry, originally as surface cultures, is widespread. Many scientists once believed that these fungi could only grow as surface cultures but it became clear in the 1940s that submerged cultures are also possible and have an enormous production potential. However, there appeared to be one problem: their form. In their natural environment filamentous fungi grow in long, branched threads called hyphae. This form, which is ideal for survival in nature, presents no problem in surface cultures, but it is often a nuisance in submerged cultures because of the strong interaction between submerged hyphae. This results in high apparent viscosities (“applesauce” behaviour) and – as a consequence – in major problems in the transport of O_2 , CO_2 , and nutrients, as well as in low productivities compared with theoretical values and with productivities obtained with other microorganisms. It was obvious that the control of the form of these fungi was a real issue that needed further attention in order to make optimal use of their potential production capacities.

Many scientists have been studying this problem from an engineering point of view for a number of decades. Simultaneously, many other scientists, working on morphology mainly because of pure scientific interest or sheer curiosity, have been very active.

The outcome of the efforts mentioned above is an impressive landscape of results about what is now called “the morphology of fungi”. This paper is about

this landscape: what it looks like, how it emerged and developed, which tools were developed, and what are its strengths and weaknesses.

2 The Framework of This Study

As will be clear from the introduction this is not another review on the morphology of fungi. There are excellent, up-to-date and extensive reviews available [1]. This is a survey of the main lines of development of a very interesting area of biotechnology research. based on a limited number of characteristic publications. These have been selected on the basis of their contributions – either good or debatable ones – to new developments in two areas:

- Improved scientific insight.
- Bioprocess practice – is it useful and usable?

The improvement of scientific insight usually goes hand in hand with a number of developments in the models used (see Fig. 1). These developments provide the main yardsticks for the present evaluation.

The trend in the development from unstructured to structured models needs an introduction. In unstructured models one assumes that the object of study has no structure: for example, a hyphal element is considered to be a more-or-less black box without internal detail. If one distinguishes septa, nuclei etc., the model then becomes structured. This structuring can go on a long way and become very detailed, but a limited number of internal “compartments” is usually sufficient to describe an observed phenomenon properly.

In the literature, models of another useful kind are sometimes mentioned: segregated – or corpuscular – models. In that case, a population is not considered to be a unit with average properties, but a collection of different individuals, each with its own properties: form, size, respiration rate, etc.

The methods used for the parameter optimization and the validation of the models will also be part of the evaluation.

Three classes of subjects will be discussed:

1. Methods: image analysis, microelectrodes, single hyphal elements, staining.
2. Models: models for morphogenesis and for the relation between morphology and production.
3. Special aspects: genetics, transport phenomena.

Now that the subjects and the yardsticks have been presented, just one word about the the author’s viewpoint. This point of view is that of a former uni-



Fig. 1. Development of models

versity professor, who started research after the morphology of moulds in 1971 and – inspired by problems he met as a consultant of Gist-brocades – worked in this particular area of biotechnology for about 10 years. After 17 years at the university, he went to Gist-brocades and worked there for 10 years. Most of the time as a director of R&D, in which position he became heavily involved with technology transfer among all of the disciplines necessary for the development of new products/processes and the improvement of existing ones.

3 Introduction to Morphology

3.1 What Is Morphology?

Morphology is the science of the form of things. It is a wide spread field of attention in a large number of sciences: biology as a whole, geology, crystallography, meteorology, chemistry – biochemistry in particular, etc. It usually starts as a way of classifying objects on the basis of their form. When the scientist becomes curious about the “why” of the development of a form he/she gets involved in the relationship between form and function. In the end, this can result in the prediction of properties given a particular form, or in the control of form/function.

First, we need several definitions. A hypha (plural: hyphae) is a single thread of a hyphal element. A hyphal element consists of a main hypha, usually with a number of branches, branches of branches etc., that originates from one spore. A flock is a loosely packed, temporary agglomerate of hyphal elements. A pellet or layer is a dense and – under normal process conditions – almost permanent configuration of hyphae or hyphal elements (see Fig. 2).

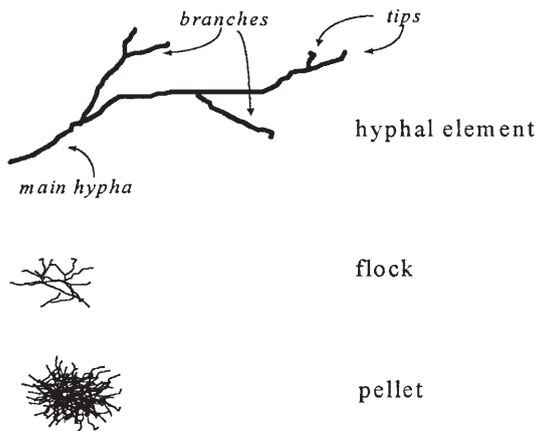


Fig. 2. Several definitions and forms

Furthermore, the “form of things” is a rather vague concept that needs further specification. The morphology of fungi is usually characterized by a limited number of variables, all related to one hyphal element: the length of the main hypha (L_e), the total length of all the hyphae (L_t), the number of tips (n) and the length of a hyphal growth unit (L_{hgu}). The L_{hgu} is defined as L_t/n .

3.2

The Morphology of Filamentous Fungi

The various forms of filamentous fungi have advantages and disadvantages in production processes as regards mass transport properties and the related overall (macro) kinetics, in particular at concentrations above 10–20 kg m^{-3} dry mass (see Table 1). As has already been mentioned, the poor transport properties are the result of the strong interaction between the single hyphal elements at high biomass concentrations, often resulting in fluids with a pronounced structure and a corresponding yield stress. This results in poor mixing in areas with low shear and in bad transport properties in general.

Morphology is strongly influenced by a number of environmental conditions, i.e. local conditions in the reactor:

1. Chemical conditions like: C_{O_2} , $C_{substrate}$, pH.
2. Physical conditions like: shear, temperature, pressure.

We will use the same notation as Nielsen and Villadsen [2] to represent all these conditions by one vector (\mathbf{z}). Thus morphology(\mathbf{z}) means that the morphology is a function of a collection of environmental conditions represented by the vector \mathbf{z} . If necessary \mathbf{z} will be specified.

Also, genetics must have a strong influence on the morphology, because the “genetic blueprint” determines how environmental conditions will influence morphology. We will return to this important issue later on. For the time being, it suffices to say that at present, despite impressive amounts of research in this area, very little is known that gives a clue to the solution of production problems due to viscosity in mould processes. This situation shows strong similarity with the following issue.

Table 1. Transport properties of various forms of moulds

Form of element	Transport to element within broth	Transport within element	Mechanical strength of element
Single hyphal elements	-/+ ^a	+	±
Flocs	-/++ ^b	±	-
Pellet/layer	+	-	+

^a Depending on the shape, size and flexibility of the hyphal element.

^b Depending on kinetics of floc formation and rupture.

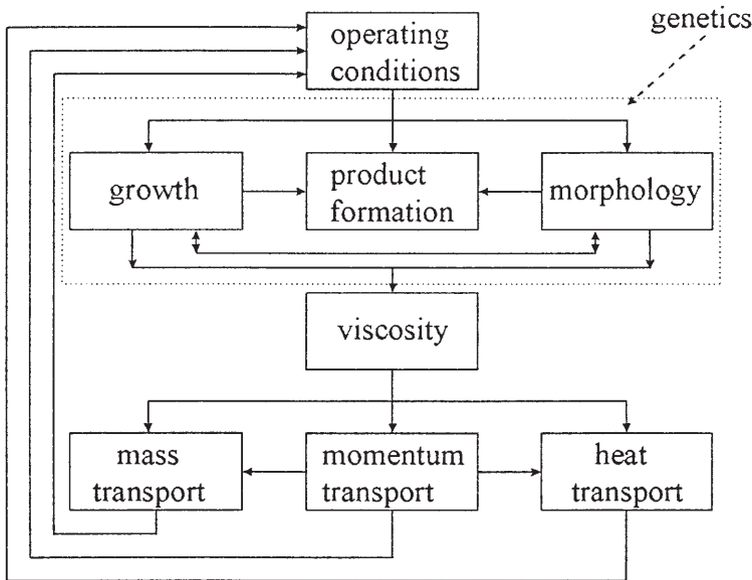


Fig. 3. Mutual influences between morphology and other properties

A very important practical aspect of the morphology of filamentous fungi is the intimate mutual relationship between morphology and a number of other aspects of the bioprocess. This has already been mentioned by Metz et al. [3], in the publication on which Fig. 3 is based. The essential difference is the inclusion of the influence of genetics. In this figure, viscosity is positioned as the central intermediate between morphology and transport phenomena. Arguments in support of a different approach are presented in Sect. 4.3.2.

This close relationship, which – apart from genetics to some extent – is without any “hierarchy”, makes it very difficult to master the process as a whole on the basis of quantitative mechanistic models. The experience of the scientists and the operators involved is still invaluable; in other words: empiricism is still flourishing.

Morphology influences product formation, not only via transport properties – as suggested by Fig. 3 – but can also exert its influence directly. Formation of products by fungi can be localized – or may be optimal – in hyphae with a specific morphology, as has been observed by Megee et al. [4], Paul and Thomas [5], Bellgardt [6] and many others.

4 Overview of the Research

This chapter comprises three topics: methods, models, aspects.

4.1

Methods

Methods are interesting because they provide an additional yardstick for measuring the development of a science. Improved methods result in better quality and/or quantity of information, e.g. more structural details, more information per unit time. This usually results in the development of new models, control systems etc. The different aspects that will be mentioned are: image analysis (Sect. 4.1.1), growth of single hyphal elements (Sect. 4.1.2), micro-electrodes (Sect. 4.1.3) and staining (Sect. 4.1.4).

4.1.1

Image Analysis

Much of the early work on morphology was of a qualitative nature. Early papers with a quantitative description of the morphology of a number of fungi under submerged, stirred, conditions have been published by Dion et al. [7] and Dion and Kaushal [8] (see Table 1 of van Suijdam and Metz [9]). A later example is the early work of Fiddy and Trinci [10], related to surface cultures and that of Prosser and Trinci [11]. Measurements were performed under a microscope, by either direct observation or photography. The work can be characterized as extremely laborious.

In their work, Metz [12] and Metz et al. [13] made use of photographs of fungi, a digitizing table and a computer for the quantitative analysis of the above-mentioned morphological properties of filamentous fungi (L_c , L_t , n and L_{hgu}) plus a few more. Although the image analysis was digitized, it was far from fully automated. Therefore, the work was still laborious, but to a lesser extent than the work of the other authors mentioned above.

The real breakthrough came when automated digital image analysis (ADIA) was developed and introduced by Adams and Thomas [14]. They showed that the speed of measurement – including all necessary actions – was greater than the digitizing table method by about a factor 5. A technician can now routinely measure 200 particles per hour. Most of the time is needed for the selection of free particles.

Since then, ADIA has been improved considerably by Paul and Thomas [15]. These improvements allow the measurement of internal structure elements, e.g. vacuoles [16], and the staining of parts of the hyphae, in order to differentiate various physiological states of the hyphae by Pons and Vivier [17].

Although the speed and accuracy of the measurements, as well as the amount of detail obtained, show an impressive increase, there are areas, e.g. models, where improvement of ADIA is essential for further exploration and implementation. An important area is the experimental verification of population balance, in which case the distribution in a population of more than 10,000 elements has to be measured routinely [18]. This is not yet possible, hampering the verification of these models. For average-property models, where only average properties have to be measured, 100 elements per sample are sufficient, and this can be done well with state-of-the-art ADIA.

Closely related to ADIA is automated sampling, which allows on-line sampling and measurement of many interesting properties, including morphology. This method is feasible but is not yet fast and accurate enough [17].

Needless to say, in all methods great care must be taken in the preparation of proper samples for the ADIA. Let this section end with a quotation from the thesis of Metz [12] (p. 37) without further comment. It reads: "The method for quantitative representation of the morphology proved to be very useful. About 60 particles per hour could be quantified. A great advantage of the method was that the dimensions of the particles were punched on paper tape, so automatic data analysis was possible".

4.1.2

Growth of Single Hyphal Elements

Measurement of the growth of single hyphal elements is important for understanding what is going on during the morphological development of mycelia. It allows careful observation, not only of the hyphae such as hyphal growth rate, rate of branching etc., but also – to some extent – of the development of microstructures inside the hyphae, such as nuclei and septa. This has contributed considerably to the development of structured models. There are early examples of this method [10], in which a number of hyphal elements fixed in a surface culture were observed. An example of present work in this area has been presented by Spohr [19]. A hyphal element was fixed with poly-L-lysine in a flow-through chamber. This allows for the measurement of the influence of substrate conditions on the kinetics of morphological change in a steady-state continuous culture with one hyphal element. This work will be mentioned again in Sect. 4.2.3.

4.1.3

Staining

Another technique that has contributed to the structuring of models is the use of staining. This has a very long history in microbiology, e.g. the Gram stain, in which cationic dyes such as safranin, methylene blue, and crystal violet were mainly used. Nowadays, new fluorescent dyes and/or immuno-labelled compounds are also being used [17, 20], allowing observation of the internal structure of the hyphae. A few examples are listed in Table 2:

Table 2. Staining

Dye	What does it show?
Neutral red	Apical segments
Methylene blue/Ziehl fuchsin	Physiological states in <i>P. chrysogenum</i>
Acridine orange (AO) <i>fluoresc.</i>	RNA/DNA (single or double stranded)
Bromodeoxyuridine (brdu) <i>fluoresc.</i>	Replicating DNA
Neutral red	Empty zones of the hyphae
Methylene blue/Ziehl fuchsin	

Applications in morphology have been mentioned [17, 20]. Several examples are:

- Distinction between dormant and germinating spores; location of regions within hyphae – as well as in pellets – with or without protein synthesis (AO).
- Propagation in hyphal elements (BrdU) in combination with fluorescent antibodies).
- These techniques contribute to the setup and validation of structured models.
- Measurement of NAD(P)H-dependent culture fluorescence, e.g. for state estimation or process pattern recognition, is also possible [21].

4.1.4

Micro-Electrodes

As has already been mentioned in Sect. 3 a (Table 1), filamentous fungi, among others, can occur as pellets or as a layer on a support. This has both advantages and disadvantages. An example of the latter is limitation of mass transfer and, therefore, a decrease in conversion rate within the pellet or layer compared with the free mycelium. The traditional chemical engineering literature had developed mathematical models for this situation long before biotechnology came into existence [22] and these models have been successfully applied by a whole generation of biotechnologists. The development of microelectrodes for oxygen [23], allowing detailed measurements of oxygen concentrations at every position within pellets or layers, opened the way to check these models. Hooijmans [24] used this technique to measure the O₂ profiles in agarose pellets containing an immobilized enzyme or bacteria. Microelectrodes have also been used to measure concentration profiles of O₂ and glucose (Cronenberg et al. [25]) as well as pH and O₂ profiles [26] in pellets of *Penicillium chrysogenum*. These measurements were combined with staining techniques (AO staining and BrdU immunoassay). This resulted in interesting conclusions regarding a number of physiological processes in the pellet.

Much of what has been mentioned above about methods, such as staining and microelectrodes, has been combined in Schügerl's review [20]. This publication also discusses a number of phenomenological aspects of the influence of environmental conditions (z), including process variables, on morphology and enzyme production in filamentous fungi, mainly *Aspergillus awamori*.

4.2

Models

4.2.1

Introduction

A majority of the models describing the morphogenesis of filamentous fungi deal with growth and fragmentation of the hyphal elements. Structured models have been used from early on. A number of them will be shown in this

paragraph, but some physiological mechanisms of cell wall formation are presented first

The basis for mechanistic, structured, mathematical models describing the influence of growth on the morphogenesis of fungi is physiology. At least, the basic assumptions of the model should not contradict the physiological facts. Therefore, a brief overview of the physiology of growth, based mainly on a publication of Gooday et al. [27], is presented here. Emphasis is on growth of Ascomycetes and Basidiomycetes, comprising *Penicillium* and *Aspergillus*, inter alia. In other fungi, the situation may be different.

Growth of fungi manifests itself as elongation – including branching – of the hyphae, comprising extension of both wall and cytoplasm with all of its structural elements: nuclei, ER, mitochondria and other organelles. The morphology of fungi is determined largely by the rigid cell wall [28]; therefore, this introduction is limited to cell-wall synthesis.

Cell-wall synthesis in hyphae is highly polarized, because it occurs almost exclusively at the very tip, the apex.

4.2.1.1

Building Blocks

The major components of the cell wall are chitin and glucan. Chitin forms microfibrils and glucan the matrix material in between them. The resulting structure is very similar to glass-fiber reinforced plastic.

Vesicles, containing precursors for cell wall components and enzymes for synthesis and transformation of wall materials, are formed at the endoplasmatic reticulum (ER), along the length of the active part of the hyphae. The concentration of vesicles in the hyphal compartment increases gradually from base to tip by about 5% by volume at the base, to 10% at the tip, with the exception of the very tip, where a rapid increase in the vesicle concentration is observed. At that point, up to 80% by volume of the cytoplasm may consist of vesicles.

4.2.1.2

Transport Mechanisms

This subject deserves some attention, because it is a common mechanism in all polarized growth models. Vesicles are transported to the tip by mechanisms that are still obscure. A number of suggestions for this transport mechanism have been summarized [27]

1. Electrophoresis due to electropotential gradients.
2. A decline in concentration of K^+ pumps towards the tip, resulting in a stationary gradient of osmotic bulk flow of liquids and vesicles to the tip.
3. A flow of water towards the tip, due to a hydrostatic pressure difference within the mycelium.
4. Cytoplasmic microtubules guiding the vesicles to the tip.
5. Microfilaments involved in intracellular movement.

Diffusion is excluded from this summary because the concentration gradient towards the tip increases (i. e. $dC_{\text{vesicles}}/dx > 0$), and therefore passive diffusion cannot play a role.

With regard to point 3, microscopically visible streaming of the cytoplasm is said to occur in fungi [29]. It is likely, however, that what has been observed is not the flow as such, but the movement of organelles. The two cannot be distinguished, because we are unable to perceive movement without visual inhomogeneities, such as particles, bubbles, clouds, etc. Moreover, the mechanism behind this movement does not have to be flow. The presence of flow is not likely, because flow needs a source and a sink. The source is present, i. e. uptake of materials through the cell membrane, but where is the sink? A sink could be withdrawal of materials needed for extension of the hyphae, but then the flow would never reach the very tip. Recirculation of the flow could be a solution for the source/sink problem but then a “pump” is needed, and it is not clear how this could be realized. Therefore transport to the apex is difficult to envisage. Passive diffusion is not possible, because the concentration gradient is positive, and flow within the cytoplasm is unlikely, because there is no sink.

An interesting hypothesis, that is an elaboration of the points 4 and 5, has recently been suggested, which can solve the problems of diffusion and flow. Howard and Aist [30] and others have shown that cytoplasmic microtubules play an important role in vesicle transport, because a reduction in the number of microtubules in *Fusarium acuminatum* inhibits vesicle transport. Regalado et al. [31] have given a possible explanation of the transport of vesicles, based on the role of microtubules and the cytoskeleton in general. They consider two transport mechanisms, diffusion and flow. In particular, their proposal for the diffusion process has a very plausible basis. In the literature, the usual driving force for transport by diffusion is a concentration gradient, but they propose a different mechanism. If stresses of a visco-elastic nature are applied to the cytoskeleton, the resulting forces are transmitted to the vesicles. The vesicles experience a force gradient that converts their random movement in the cytoskeleton to a biased one. Consequently, they move from regions of high stress to regions of low stress. The driving force is thus no longer a concentration gradient but a stress gradient, thus solving the problem that arose with the classical diffusion. Their computer simulations look very convincing, but more experimental evidence is needed. They cite many other examples from the literature showing the relation between cytoskeletal components and vesicle transport.

4.2.1.3

Synthesis of the Cell Wall: Chitin

Chitin synthesis occurs exclusively at the growing hyphal tip and wherever cross-walls (septa) are formed. This indicates that chitin synthesis has to be closely regulated both in space and time. All of the genes for chitin synthase that have been isolated so far code for a protein with an N-terminal signal sequence. This indicates that the protein is synthesized at the ER, transported through the Golgi and brought to the site of action, i. e. the hyphal tip or the site of cross-

wall formation, in secretory vesicles. There, it functions as a transmembrane protein, accepting the precursor UDP-N-acetyl-glucosamine at the cytoplasmic site and producing chitin polymers on the outside. There, in the wall area, different chitin polymers interact by mutual H-bonding, and crystallize spontaneously into microfibrils. This crystallization process might be hampered by the cross-linking of newly synthesized chitin to other wall components.

Early findings for many fungal chitin synthases were that these enzymes are often isolated in an inactive state and can be activated by proteolytic digestion [32]. This led to the idea that synthetases may be regulated by transformation of a zymogen form into an active enzyme, but not necessarily by proteolysis. Treatment of membrane preparations with detergent resulted in loss of activity that could be restored by addition of certain phospholipids, indicating that the lipid environment in the membrane might be another possible activating factor.

4.2.1.4

Synthesis of the Cell Wall: Glucan

The enzyme involved in glucan synthesis is also a membrane bound protein that catalyses the transfer of glucosyl residues from UDP-glucose to a growing chain of β -1,3- linked glucosyl residues. It was found that the synthase is highly stimulated by micromolecular concentrations of GTP. Subsequently, testing mutants for GTP-binding proteins with a phenotype compatible with a defect in cell wall synthesis, *Rho 1*-mutants were found. Tests of these mutants established unequivocally that Rho 1 protein (Rho 1p) is the GTP-binding protein that regulates β -1,3-glucan synthase and is essential for its activity [33].

Intriguingly, Rho 1p has two further functions: it regulates both cell-wall synthesis and morphogenesis. Rho 1p activates protein kinase C, which in turn regulates a pathway that leads to cell-wall synthesis in response to osmotic shock [34]. However, Rho 1p is not directly involved in the activation of β -1,3-glucan synthesis. Furthermore, Rho 1p may also be involved in the organization of the actin cytoskeleton at the hyphal tip (Yamochi et al. [35]).

4.2.1.5

Synthesis of the Cell Wall: the Structure

Finally, covalent bonds are formed between chitin and glucan polymers, and hydrogen bonds are formed between the homologous polymer chains, resulting in a strong combination of chitin fibers in a glucan matrix (Wessels [36]). Wessels also makes it clear that the spatial distribution of wall extrusion and progressive crosslinking might result in different morphologies: mycelium, pseudo-mycelium, and yeast.

There are additional compounds present in the membrane and other enzymes are also involved, but the essentials needed for the evaluation of the growth models in this paragraph have been presented above.

It should be clear that the growth of the cell wall is a rather complicated process, with many enzymes involved and much that is still unknown, but a number of facts are known that can exclude certain mechanisms.

Before we start the discussion about mathematical growth models, two more general remarks regarding structured models have to be made. Structured models are not necessarily of a mechanistic nature. One can observe one hyphal element under the microscope and describe all the internal structures one sees exactly, without any mechanistic explanation. This description can also be quantitative but it remains empirical (*the “flora” is a typical example of a non-mechanistic structured approach: it shows all the different organisms in a habitat – and often their parts as well – in a systematic way, without explaining “why”*).

The models that we will discuss have been validated by experiments with various fungi. However, the morphological characteristics of moulds vary enormously between different strains, even between strains belonging to the same species. In other words, the quantitative results are very specific. Therefore, the main values of a well-validated model are the methodology and the structure, not the actual figures.

4.2.2

Morphology Modelling in General

A systematic survey of the modelling of the mycelium morphology of *Penicillium* species in submerged cultures has recently been published [18]. The authors distinguish between various kinds of models (see Fig. 4). A short explanation follows.

Models of Single Hyphal Element. Experiments with single hyphal elements were mentioned in Sect. 4.1.2. The advantages, greater detail and more insight, can be used in single hyphal element models. Several examples have been mentioned [11, 18].

Population Models. In population models, or population balances, microorganisms are treated as individuals with different properties. Each individual hyphal element has its own properties, in this case usually L_t and n . Central in these models is the population density function, $f(L_t, n, t)$, representing the number of individuals per volume in the population with a specific value of L_t and n at time t . The value of $f(L_t, n, t)$ can change under the influence of tip

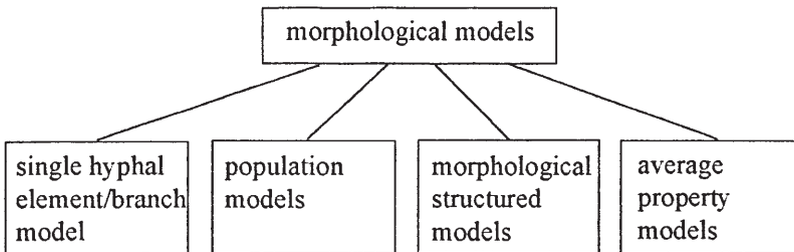


Fig. 4. Kinds of morphological models

extension, branching, birth of new hyphal elements due to germination or fragmentation, and – in continuous cultures – dilution.

Population balances were used in the biotechnology of bacteria and yeasts, before the term had been coined (e.g. [37]), but not for filamentous fungi. The main reason is that the verification of these models required rapid methods for measuring the properties of the individuals. This is no problem for individuals of the size of bacteria and yeasts. Thousands of cells can be measured quickly and routinely (Coulter counter). For filamentous fungi the situation is different. For the characterization of the morphology of fungi, not one but at least two variables are important, e.g. L_t and n . As mentioned in Sect. 4.1.1, they have to be measured for so many elements that it is too much even for ADIA at present. Therefore, we will not deal with these models.

Nevertheless, these models are very elegant and allow for very detailed description of microbial systems [18]. One thing is certain, the processing and storage capacity of computers will increase drastically so that the prospects for use of these models in the future are favourable.

For those interested in the background of population balances, two publications are recommended: Randolph [38] and Randolph and Larssen [39].

Morphologically Structured Models. These models deal with conversions between different morphological forms, resulting in shifts in fractional concentrations of biomass with a specific morphological form (Nielsen [40]).

Average Property Models. These models deal with the averages of the population as a whole, i.e. average length, average number of tips per hyphal element (e.g. [12, Aynsley et al. 41, Bergter 42]). This group forms the vast majority of all models – not only in morphogenesis!

We will deal with models for the development of a particular morphology (morphogenesis) and models that include the influence of the morphology on the production of metabolites – usually antibiotics.

4.2.3

Models for Morphogenesis

These models contain only mechanisms for growth, branching and fragmentation. In fact, these “single” mechanisms are usually the result of underlying submechanisms, and so on. One well-known example has already been mentioned [11]. The only mechanism is growth but, as we will see below, there are seven or eight submechanisms, depending on the degree of subdivision.

Morphogenesis is the development of a particular morphological form. For fungi this form is usually characterized by the number of tips and either the total hyphal length (L_t) or the length of the main hypha (L_e) (both per hyphal element).

The first models dealt only with growth. Examples are the constant linear extension rate of pellets and hyphal colonies, apart from the very beginning of hyphal growth (Trinci [43, 44]), the branching after a certain length of the hypha (the first structuring (Plomley [45])), and the constant value of the

L_{hgu} (Caldwell and Trinci [46]). The L_{hgu} is not constant if the diameter of the hypha varies [40]. Much more constant is the mass of a hyphal growth unit (m_{hgu}) where $m_{\text{hgu}} = m/n_{\text{tot}}$.

Because branches also grow at a constant rate, form new branches etc., this results in exponential growth of the mycelium under non-limiting substrate conditions.

One of the other older growth models worth mentioning is the cube-root law for the growth of pellets (Emerson [47]). It was presented as an empirical model, but the mechanism behind it is obvious once one realizes that, under the condition of substrate limitation, growth of a pellet can only occur in a thin layer of almost constant thickness. The amount of structuring in these models is rather low or absent.

The study of single hyphal elements [19] revealed that the tip extension rate of the hypha can be modelled with saturation kinetics with respect to the branch length. Nielsen and Krabben [48] found the same relation for the average tip-extension rate when samples of about 50 hyphal elements were measured.

The next group of models are representatives of morphogenesis as a result of growth and fragmentation.

The earliest work on fragmentation was performed at the Istituto Superiore di Sanità in Rome [7, 8] where extensive phenomenological research has been done on the influence of stirring on the morphology of *Penicillium chrysogenum* and a number of other filamentous fungi. They noticed a strong correlation between morphogenesis and stirrer speed, but did not correlate their results in mathematical form. This was done by others [12] and has also been published in a review [9]. The dependence of the effective length (L_e) on P/V could be represented by $L_e = \text{const.} \cdot \epsilon^\alpha$, where ϵ represents the power per unit mass of the broth. The value of α varies from -0.66 to -1.11 .

Growth and fragmentation of *Penicillium chrysogenum* were combined in one model rather early [12, 9]. The growth model used was based on early work [45, 46]. Hyphae grow at a constant linear rate. After $\Delta L = L_{\text{hgu}}$, a new branch is formed, etc. This results in exponential growth of the total length per hyphal element, and of the culture as a whole. The only structure included in the growth model is branching. The model used for fragmentation is based on the turbulence theory of Kolmogorov (see [49]). Breakup of the main hypha is due to eddies formed as a result of local energy input into the medium. First, the maximum effective length of the hyphal element $L_{e_{\text{max}}}$ before fragmentation occurs is calculated. $L_{e_{\text{max}}}$ is the maximum length of the main hypha):

$$L_{e_{\text{max}}} = \text{const.} \cdot \frac{d_h^{0.38}}{\epsilon_{\text{max}}^{0.25}} \quad (1)$$

The constant includes, inter alia, wall thickness and tensile strength of hyphae; d = diameter of hyphae.

The main hypha will break up only if $L_e > L_{e_{\text{max}}}$. This will only occur in a limited volume of the reactor near the stirrer (V_{disp}), where ϵ is at a maximum.

The dynamics of the fragmentation process is introduced by assuming that its rate is determined by the probability of fragmentation, which is proportional to the number of times the element passes V_{disp} .

V_{disp} is a function of ϵ , N , D_r and the distance to the stirrer (r). The final result is:

$$-\frac{dL_e}{dt} = \text{const.} \cdot N^{1.75} \cdot D_R^{0.5} \cdot d^{-0.38} \cdot (L_e^2 - L_{e_{\max}}^2) \quad (2)$$

for $L_e \geq L_{e_{\max}}$, otherwise $dL_e/dt = 0$.

This is one of the first detailed mechanistic models for the fragmentation of hyphal elements. Among other factors, it takes into account the fact that the turbulent energy is absorbed in a small region near the stirrer, so cannot be considered to be evenly distributed. As we can see, dL_e/dt is proportional to L_e^2 . Other authors assume a different dependence on L_e .

The steady-state value of L_e can be calculated assuming that the growth rate is equal to the fragmentation rate. The approximate result is:

$$L_e = k_1 d^{0.38} / N^{0.75} D_R^{0.5} + k_2 \mu / N \quad (3)$$

The growth and fragmentation of pellets has been studied by van Suijdam [50]. The model of van Suijdam includes growth and autolysis, mass transfer of oxygen – described as density-dependent diffusion accompanied by chemical reaction, as well as external oxygen transfer from bubble to liquid and from liquid to pellet. The basis for the fragmentation is again the Kolmogorov theory [51]. The mechanistic model for fragmentation of the pellets can be summarized as follows: $d_{\text{aver}} = c\epsilon^{-0.4}$, where c is a constant; the experimentally determined power of ϵ was 0.38.

To obtain a model for spore germination, growth and fragmentation of *Penicillium chrysogenum* a population balance was used as a start to derive balances for average properties [48]. By using the results of their own experiments and experimental data from the literature, it was possible to extract interesting information about germination, growth and fragmentation of hyphal elements. A very good fit was obtained between model and experiments for those periods during the bioprocess in which the conditions for the extraction were fulfilled, e.g. no fragmentation during germination.

Ayazi Shamlou et al. [52] present a fragmentation model with first-order dependence in L_e :

$$\frac{dL_e}{dt} = k \cdot (L_{e_{\text{equil}}} - L_e) \quad (4)$$

They too use the Kolmogorov theory to calculate $L_{e_{\text{equil}}}$ but they assume that the turbulence in the vessel is homogeneous and isotropic, which is not correct.

The models mentioned above have a very limited physiological background. This is particularly evident in the case of the fragmentation models, although there certainly are opportunities to include physiological mechanisms in fragmentation, for example, vacuolization and subsequent decrease in the local

tensile strength of the hyphae. The growth models are also almost non-physiology based; only L_{hgu} has a physiological background.

We will now deal with a number of physiology-based models.

An often quoted example of an early mechanistic and structured growth model of a single hyphal element, based on solid physiological observations [11], is an extension of earlier work of the same group [10]. The structural elements are: tips, septa and the regions between them, vesicles and nuclei. Growth – in fact, cell-wall synthesis – occurs at the tips of the hyphae as a result of the *inclusion of vesicles*. These *vesicles are produced* at a constant rate at the wall of the hyphae, *transported* to the tips and used for growth. Under the influence of growth, the ratio of cytoplasmic volume and the number of nuclei at the tips in the apical compartment attains a critical value. This triggers *an increase in the number of nuclei* until they have doubled (from four to eight). These eight nuclei are then divided into two sections, each of four nuclei, by *the formation of a new septum* between them, and so on. These septa have an *opening, the diameter decreases with age*, i.e. with distance from the tip. This results in a *congestion of vesicles* before such a septum, which – in turn – triggers *branch formation*. The submechanisms have been *underscored*.

Several other attempts have been made to increase the physiological basis, or the structuring, of the growth mechanism in models. An example [41] is the use as a growth model of a self-extending tubular reactor that absorbs nutrients along its length. One of the assumptions is transport of the precursor through the hyphae at a constant average rate of flow. As has already been mentioned in Sect. 4.2.1, this is not a very likely mechanism, either from a physical or from a physiological point of view. Furthermore, a number of empirical kinetic equations have been used, to which four of the seven parameters in the model have been fitted.

It is very interesting to compare three models. When the hyphal element continues to grow, a pellet can emerge. Yang et al. [53] and King [54]), working with *Streptomyces* – which is not a fungus but a member of the prokaryotes – and Bellgardt [6], working with *Penicillium*, start with the growth of hyphae but deal mainly with the growth of a pellet from a single spore. These models show a considerable amount of structure; they contain submodels for growth, septation and branching of hyphae, and simulate the observed phenomena very well. They show quite a few family features but clear differences as well.

The part of all these models that deals with hyphal growth is based on a model that combines diffusion with chemical reaction (DCR). The chemical reaction concerns the production – in one model the degradation as well – of a key precursor of growth. Production is either constant or contains a saturation term. Degradation is first order. For septation, a Trinci-like approach was used. The models contain several stochastic elements: branching occurs near the septa but is normally distributed, and the angles for branching and growth are normally distributed as well.

DCR models have also been used to calculate oxygen and other substrate profiles in the pellet. King [54] uses an purely kinetic equation for the elongation and fragmentation of hyphae and a DCR equation for the formation and fragmentation of tips in the pellet. In the model of Bellgardt [6], mycelial

growth in the pellet is described the same way as hyphal growth. The diffusion coefficient in the DCR model for the substrate depends on the local density of biomass, and therefore on its radial position in the pellet. This model contains a term for the fragmentation of tips that grow out of the dense part of the pellet. This fragmentation was described with a probability function. Fragmentation of the hyphae occurs only at the tips.

Not too many parameters are involved in this modelling, and a number of them can be measured separately a priori.

The assumptions underlying these models are shown in Table 3. Table 3 shows how a limited number of basic mechanisms can result in rather differently structured models. The amount of structuring can still be increased, but at a cost as will be discussed in Sect. 4.2.5.

Table 3. Comparison of a number of highly structured models

	King	Bellgardt	Yang et al.
Hyphae			
Hyphal elongation eq.	DCR ^a	DCR	DCR
Hyphal elongation kinetics	Zero-order production 1st order degradation of key component	Zero-order production	Logistic production ^b
Transport in hypha	Passive diffusion	Passive diffusion	Passive diffusion
DNA replic.	Logistic production	n.i.	Logistic production
Septum formation	Deterministic	n.i.	Deterministic
Branching	Stochastic	Stochastic	Stochastic
Pellets			
Growth	From one spore	From one spore	From one spore
Hyphal elongation eq.	Kinetics only (no mass transfer)	See hyphae (above)	See hyphae (above)
Kind of kinetics	growth, degradation, fragmentation	See hyphae (above)	See hyphae (above)
Tip formation	DCR	Stochastic	Stochastic
Tip formation kinetics	Tip formation degradation, fragment	n.r.	n.r.
Substrate consumption	DCR	DCR	n.i.
Density	Growth dependent	$f(r_{crit})$	n.i.
Diff. Coef.	Based on void volume	$f(\text{cell density})$	n.r.
Fragmentation of tips	Included in kinetics	Probability function	n.i.

n.i., not included; n.r., not relevant.

^a DCR: an equation containing terms for diffusion and (chemical) reaction kinetics.

^b Logistic equations are of the form: $r(C) = k \cdot (C_{max} - C)$. There is also another definition of a logistic equation! [55].

As has already been mentioned, experimental confirmation of the models is not bad at all. This does not mean, however, that the models are correct from a physiological, or even physical, point of view (see Sect. 4.2.5). It is clear from Sect 4.2.1 that classical diffusion makes no sense. Yang [53] assumes explicitly that there is no active transport of the key component in the hyphae and zero concentration of the key component for hyphal elongation at the tip. This can only be true if the vesicles do not contain the key components. Furthermore, it is obvious that the kind of kinetics used for hyphal extension describes what is going on quite well, but its choice seems rather arbitrary. Interestingly enough, it does not seem to be important whether or not the production term for hyphal elongation is zero order or logistic, and whether a first order degradation term has been included. This raises an interesting, not merely philosophical, question: What do we mean when we say that a model is mechanistic?

To conclude this section, an impressive quantity of work has been published about the influence of environmental factors ($z(t)$) on morphology. These results are mainly of an empirical nature, although suggested mechanisms are not uncommon. However, structured mechanistic models in this area, based on intracellular reactions at a molecular level, are virtually absent. No examples are presented here.

4.2.4

Models Describing the Relation Between Morphology and Production

Fungi are important workhorses in the fermentation industry, and models that allow for optimization of production with fungi can be very useful. In a special group of models morphology has been related to production. If the production of a component is dependent on the morphology of a microorganism as such, rather than via the influence of morphology on bulk transport, the use of morphologically structured models is a must. A number of models have been set up to describe this relationship, with the final goal of optimizing productivity. These models generally show a high degree of structure, and describe the production process quite well, at least at laboratory scale.

Three examples will be presented: first, an old one [4], followed by two recent ones [5, 6].

The first model [4] contains five “differentiation states”, four of which are connected with a particular product, whereas the fifth is the active growing tip. The concepts of the model are suggested by data from experiments of the authors with *Aspergillus awamori*. The different morphological forms have been observed on surface cultures. A highly structured model was set up for batch and continuous cultures. The 43 parameters were obtained from the literature and their own work. The phenomena observed during the simulation find a number of parallels in the literature, but there is no direct comparison between their simulations and detailed data from parallel experiments. An extensive overview of this model has been given by Nielsen [56].

In a morphologically structured model for production of cephalosporin with *Streptomyces* on complex substrates [6], four different morphologies were discerned:

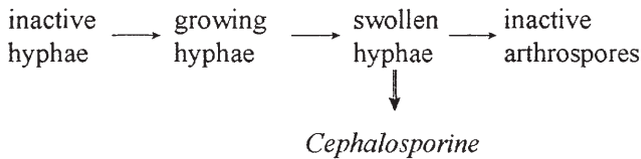


Fig. 5. The four morphological forms of *Acremonium chrysogenum*

The model consists of balances among these morphological forms, and a number of kinetic equations for growth and product formation and for substrate consumption with Michaelis–Menten-like substrate dependence. Repression of oil consumption in the presence of sugar is also included.

The model reproduces quite well the experimental time dependence of the concentrations of cell mass, oil and sugar substrate, product, and CO₂ in the off gas, also for several cultures. The model contains 20 parameters. It is not clear from this publication how these parameters were obtained. The model is applied to the dynamic optimization of feeding strategies.

If there are no morphological differences between hyphal elements, it is still possible to look at regions in hyphae with different activities [5]. Five different regions are distinguished in hyphae of *Penicillium chrysogenum*, based on different structures and activities. The model contains 20 parameters. Eleven parameters were obtained from the literature, 9 parameters were obtained from a continuous reference process. The fit between the simulation with the model and the data of this experiment was very good, but that is not surprising with nine parameters to tune the model to the experiments. However, the same model, with the original parameters obtained from the continuous culture, was used to simulate the outcome of a fed-batch culture. Again, the results were very good. This increases the value of the model. The sensitivity of the model to the various parameters used was also checked by means of a very simple – but very useful – technique for finding out which parameters are really important and need to be known with high accuracy and for which parameters evaluation to within an order of magnitude is sufficient. This technique has also been used by others [12].

4.2.5

Some General Remarks About Models

This is the place to make a few remarks about the models we saw and about models in general. A model is always a simplification of reality. It is, as someone once said, the art of balancing between unwieldy complexity against oversimplification. First, one quotation to “set the scene”:

It should be noted, however, that a simulation giving realistic pictures does not necessarily prove the correctness of the assumptions used [53].

Almost 20 years earlier Topiwala [57] made a similar remark. In fact, models need be neither mechanistic nor structured in order to give a good correlation between model and experiment. Whether or not we have to worry about this

depends on the objective of the model, in particular on whether this objective requires some form of extrapolation from the model, although even interpolation can be a risk. Important objectives are:

- Process control: Any model – including non-mechanistic models like neural networks – will do, as long as the process is not extrapolated beyond the range of conditions in which the model has been tested.
- Process optimization: The objectives are the same as for process control, but unreliable if the optimum lies outside the test range, i.e. requires extrapolation.
- Scale-up: Scale-up is extrapolation *par excellence*, and is therefore very unwise.
- Satisfaction of curiosity: Here, the goal is to unravel the real mechanisms, in order to construct a model that is solidly rooted in physiology and physics. Jeopardizing the model, by extrapolation, *inter alia*, is the essence of the game; or “We should ring the bells of victory every time a theory has been refuted” (Popper [58]).

The objective determines a number of other issues that call for explicit decisions.

Time Dependence vs (Quasi) Steady State. This is a rather obvious decision to be taken. Time dependence can result in rather awkward differential equations, but very often a quasi-steady state approach yields acceptable results. For example, the concentration profile of O_2 in a growing pellet can be calculated very accurately without taking into account its momentary growth.

Number of Compartments. A compartment can be a geometrically defined compartment, an event, e.g. hyphal tip extension or septation, or a number of morphological forms – each with substrate consumption, product formation etc.

Mechanisms per Compartment. These mechanisms include the following:

- Transport: flow, diffusion (ordinary or active), pushing aside.
- Kinetics: nth order, logistic, saturation (e.g. Michaelis–Menten etc.).
- Equilibrium: thermodynamic, mechanical, saturation (full is full).

The choice of the mechanisms involved in each compartment is often rather arbitrary and not well argued. As stated above, this is not necessarily a problem. However, if the model has to be used for extrapolation or if the driving force is curiosity, full attention must be paid to this aspect. Some other aspects of the choice of mechanisms are as follows.

A number of mechanisms can be chosen, among others diffusion and flow, for the transport of vesicles to the tip of hyphae. These two mechanisms are compared in the Appendix. At first sight, both mechanisms give equally satisfactory results. Both are impossible, however. In order to see why, one has to look at $(C_{\text{vesicle}}(x))$ in greater detail.

Equations (8) and (10) in the Appendix can also be written as $C_1 = c_1(L^2 - x^2)$ and $C_2 = c_2x$, respectively, where c_1 and c_2 are constants. This “lumping of

parameters” facilitates parameter optimization, but makes it difficult to jeopardize the model, because one loses sight of the constituent mechanisms.

Considering the enormous concentration of vesicles at the very tip of the apex, one might consider the possibility that something other than transport of vesicles, perhaps the inclusion of vesicles in the membrane, is rate-determining.

A rather extensive set of physiological events contribute to hyphal tip extension (see Sect. 4.2.1). In most publications, this boils down to a simple Michaelis–Menten kinetic equation for one component. Maybe this is correct, but it is seldom well argued.

As mentioned above, a good fit between model and experiment may be quite independent of the submechanisms, but it usually has a great deal to do with the next point.

Parameter Estimation. For extensively structured models, parameter estimation is often more time-consuming than the setup of the model itself. The problem is the number of parameters required by the model: roughly five parameters are needed per “compartment” If the compartment is geometrically defined, there are as many as three geometric parameters, depending on its shape; the diffusion coefficient and/or linear velocity; and at least two kinetic parameters. Thus, a model with four well-defined compartments needs 20 parameters [5] and one with three phenomena, e.g. hyphal extension, septation, and branching, needs 13, including those required for specifying initial conditions [53]. The important question is: “Where do these parameters come from?” Are they from “first principles”, from the literature, independent measurements carried out separately *in vitro* and *in vivo*? Most important of all: How many parameters have been used to “fit” the model to experiment?

First, modelling the behaviour of a system and using the behaviour of the same system to determine the parameters is a form of inbreeding. As mentioned in the section on the choice of mechanisms, this is no problem if the model is to be applied in the same range of conditions as was used for determining the parameters.

If use of the model includes extrapolation, the situation is different. One should then jeopardize the model by working with the fungus under other environmental conditions ($z(t)$), other process modes, e.g. batch, fed-batch or continuous, and/or other dilution rates or stirrer speeds. The agreement between the experimental data and the predictions of the model, without further adaptation of the parameters, is a good measure of its robustness.

Second, any system can be “modelled” with 20 or 30 parameters. An old, but still very relevant, publication in this respect has the intriguing title, “The least squares fitting of an elephant” [59].

The almost exponentially increasing performance/price ratio of computer systems makes parameter fitting easier and easier. Unfortunately, this is often at the expense of observations, because the performance/price ratio of modern analytical apparatus remains much more constant.

An often neglected problem is the effect of scale-up on the values of parameters. The morphology is very dependent on environmental conditions ($z(t)$), which are themselves very often dependent on scale. In particular, a change of

regime, for example, from kinetic to transport control, can be very misleading. The vast majority of parameters have been obtained from small-scale experiments, because production scale parameters are not usually available for publication.

The substrates on the laboratory scale and on the production scale often are – and have to be – quite different [60]. This also affects $z(t)$.

The number of inoculation steps increases from 1 on the laboratory scale to about 4 on production scale. This too can influence the behaviour of the fungus.

These are typical scale-down problems: how to create a small-scale environment that is representative of the full-scale conditions.

In conclusion, the physiological basis of many models, including those in present use, has to a large extent been one publication [11], together with a small number of related publications. This publication has been exploited extensively, and for very good reasons, but we now need to make further advances. Today, tools are available for elucidation of the biochemical and genetic background of the mechanisms involved, although the task will not be an easy one. Close collaboration between those who construct models of the types mentioned above and mycological physiologists and geneticists would be a great help. A serious problem is that mycological physiologists are relatively scarce. In the Netherlands, for example, fewer than one in ten microbial physiologists is a mycologist and the ratio is probably not much better elsewhere.

4.3

Special Aspects

So far only methods and models have been presented that primarily allow for a better description and understanding of the morphogenesis of filamentous fungi. They also give some clues for the improvement of morphology. However, there are more methods for the improvement of the morphology (see Table 4 for an impression of the methods that could be used).

Some important issues can be extracted from Table 4 about how morphology can be influenced, but we will deal exclusively with genetics, because at present the other methods are obtained almost exclusively from measurements based on empirical know-how.

Furthermore, insight into the relation between morphology and transport phenomena in a mycelial reaction mixture can be of some help for the solution of morphology-based problems in production processes.

Table 4. Methods of influencing morphology

Genetics (either r-DNA or CSI)	Other methods
1. Primary cell wall synthesis	1. Inoculation
2. Altered metabolism	2. $z(t)$ (including shear rate)

CSI is classical strain improvement.

4.3.1

Genetics

Morphology is a phenotypic property. The development of a phenotype, e.g., a fungus with a specific morphology, is always the result of the genotype and the environment, a relationship that can be formulated: Phenotype = Genotype \times Environment, i.e. $P = G \times E$. As mentioned in Sect. 4.2.3, there is an impressive quantity of literature, albeit fragmented, about the influence of environmental factors on morphology; as has been mentioned above, it is mainly of an empirical nature. However, very little knowledge, even empirical knowledge about the effect of genes on morphology is available. This is a real stumbling block, for if we want to master morphology we cannot do without genetics.

The genetics of morphogenesis can be very complicated. A non-fungus example of the effect of genes on morphology is the endospore formation of a "simple" bacterium: *Bacillus subtilis* [61]. At least 45 separate loci involved in endospore formation have been identified. Some consist of single genes, others consist of gene clusters. Also, the discussion of genetics in the introduction to physiology (Sect. 4.2) shows the complexity of the genetics and its control.

Morphology is determined to a large extent by the structure of the rigid cell wall. This means that mutations that affect this structure also affect morphology. However, other processes in the cell can also affect morphology and are candidates for consideration as factors that can be influenced by genetics. Although there is little hope for the elucidation of the detailed genetics behind morphogenesis of fungi in the near future, there are several leads to possible shortcuts. For example, changing a single gene can have a drastic effect on morphology.

1. Brody and Tatum [63] demonstrated that a point mutation in one gene, resulting in a change of affinity of a single enzyme (glucose-6-P dehydrogenase) to its substrate, effected a drastic change in the morphology of a surface culture of *Neurospora crassa*, probably because of in vivo accumulation of glucose-6-P. Furthermore, the change in the affinity of the enzyme for its substrate was caused by the alteration of a single amino acid in its structure. It is also interesting that changes in the source of carbon now had no obvious effect on the newly produced morphology. It has also been mentioned that the morphology of *Neurospora crassa* can be changed by altering the genes effecting cell wall synthesis [28].
2. Nonaka [62] mentions an interesting example of the effect of small genetic changes on the morphology of *Saccharomyces cerevisiae*. He investigated the effect of the protein Rho1p on the budding of *Saccharomyces cerevisiae*. Rho1p switches glucan synthesis on at budding and off at maturation of the daughter cell. Rho1-depleted mutants stop growing with small budded cells.

The experimental results with *Neurospora crassa* [63] and with *Saccharomyces cerevisiae* [62] show that changing one gene can have a dramatic effect on morphology. If these methods could be applied in some way to other fungi as well, the use of enrichment cultures would be an interesting option. This method is, of course, almost 100% empirical, but it may lead to a rapid im-

provement in the morphology of strains, and it may help find out what molecular biologists have to look for, and where. With some creativity, it should not be too difficult to set up a system that selects for the preferred morphology. This selection may take some time, but it may well pay off. The selection of a special form of microorganism (pellets), necessary for the large-scale anaerobic treatment of wastewater, took more than a year [64].

The overall effect must be a stable increase in productivity in the full-scale reactor, as a result of the combined positive effects of the genetics on the inherent productivity of the fungus as such, as well as on the transport properties, due to its effect on the improved morphology.

4.3.2

Whole Broth Properties

As seen in Fig. 3, there is an intricate relationship between viscosity and transport properties. Furthermore, it is well known that morphology has a strong influence on viscosity [9, 12]. Because poor transport properties can ruin full-scale productivity, these relations are very important (see Fig. 6, which is a variation on the theme of Fig. 2).

It appears to be difficult to represent the morphology of fungi unequivocally by rheological equations, although many attempts have been made by Metz [9, 12] and Olsvik [65] and their respective co-workers, among others. The next difficult step is finding the relation between the often rather peculiar viscosity equations and transport properties. The last step, going from transport properties to productivity, is an application of basic chemical engineering. This is route A in Fig. 6.

One can also save a lot of trouble and choose the more phenomenological approach of route B in Fig. 6, which calls for finding a direct relation between morphology and transport properties.

The reason for the difficulties of route A are threefold. First, the consistency – a better word than viscosity – of the filamentous mixture originates

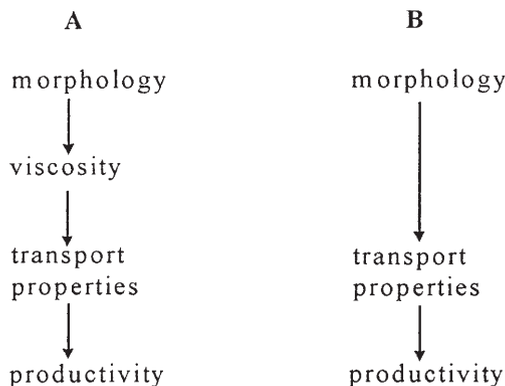


Fig. 6. From morphology to productivity

almost entirely from the interaction between solid particles. Second, these particles have almost macro-dimensions: from 0.1 mm for hyphae to several mm for flocks and pellets. Third, the consistency appears to be time-dependent.

This suggests that a dynamic corpuscular approach would be more successful than a static continuum approach. The latter is what one usually finds in the literature.

Some first steps have been made along route B [9]. Dynamic flocculation/deflocculation processes appear to be taking place in stirred mould suspensions. Deflocculation proceeds very rapidly under high-shear conditions, but in quiescent regions complete flocculation also occurs within about 0.5 s. This process time is only slightly dependent on dry mass-concentrations (C_X) in the range between 13 and 20 kg m⁻³, but increases to $\cong 5$ min for $C_X < 0.1$ kg m⁻³. Although the flocculation/deflocculation process is very fast, a number of interesting phenomena can be observed in bubble columns.

When bubbles rise in a flat plate bubble column, the wake of a bubble has an appearance that is different from the bulk of the broth. This effect may be the result of demixing alone or of demixing with its subsequent influence, due to the lower local value of X on the breaking and reforming of the structure of the broth. In any case, if successive bubbles are released, one can see that the first bubble travels much more slowly than the second. This second bubble obviously profits from the path made by the first, and so on. The overall effect is a very rapid coalescence of bubbles, growing from 5 mm to 5 cm in diameter within a 1-m column. In some full-scale mycelial processes, with a very interactive mycelium at high biomass concentrations, this results in the aforementioned applesauce appearance of the reaction mixture and football-sized bubbles, bubbles that would be too unstable to ever exist in a homogeneous fluid. Mass transfer would be extremely slow if it were completely dependent on these bubbles.

Mass transfer in a thick mycelial reaction mixture is in practice not as bad a mass transfer mechanism as might be expected. It has therefore been suggested [9] that mass transfer is not based on the contribution of bubbles, but on the intense breakup and oxygen saturation of flocs near the stirrer. The resulting equation has the following form:

$$k_1 a = \frac{\Phi_p}{V} = \frac{1}{t_c} \cong \frac{N \cdot D^3}{V} \quad (5)$$

and gives the right order of magnitude for $k_1 a$ in full-scale reactions.

Another aspect of the structure of the broth is the entrapment of small bubbles in the mycelial network. This acts as a sink or source for O₂ and CO₂.

This kind of modelling has obviously not generated many followers.

Problems with the measurement of viscosity of filamentous reaction mixtures are also the result of the inhomogeneous character, as was mentioned above.

The measurement of rheology has improved since the first measurements with a turbine impeller in the laminar flow region [9, 12, 66]. Experiments with

classical rotating cylinders for rheology measurements were unsatisfactory for three reasons:

1. The particles are of the same order of magnitude as the annulus, resulting in damage of pellets and flocs during measurement.
2. Less dense layers are formed near the wall.
3. The particles have a tendency to settle.

Tube viscometers avoid problem 1, but need a relatively large diameter when pellets as large as 1 mm are used, in which case large samples are needed [66].

Allan and Robinson [67] have shown, with suspensions of three different moulds, that a helical ribbon is preferable to the turbine impeller for viscosity measurements, because the assumption that the average shear rate in the turbine viscometer is independent of fluid rheology is not correct – at least not for fluids with a low flow-behaviour index (n) in the power-law equation:

$$\tau = K \cdot \dot{\gamma}^n \quad (6)$$

and for fluids with a yield stress. Their rotating cylinder viscometers and pipes with $D > 10$ mm worked well, contrary to earlier experience [66]. This could be because different strains and suspensions without pellets were used [67].

An interesting issue is the relation between the rheological properties and the morphology of the hyphal clumps. A positive correlation was found between the roughness of these clumps, i.e. aggregates of micelium, and the biomass concentration of *Aspergillus niger*, on the one hand, and the consistency index K of the power law equation, on the other [65]. This correlation holds for various values of the growth rate and the dissolved oxygen tension (DOT) and for two different values of the biomass concentration.

5 Implementation of the Results

The amount of research carried out in the past on the morphology of filamentous fungi has been impressive. However, the implementation of these results in the fermentation industry seems to be limited. There are a number of reasons for this limited implementation:

The first reason is the problem of time squeeze. In the fermentation industry, one is constantly looking for more productive strains. Per product, this results in about one new, more productive strain every few years. This strain is taken into production as soon as possible, and any morphology problems are solved by empirical methods based on extensive experience. Schügerl [20] points out the complex interrelationships between all the factors involved in growth, morphology, physiology and productivity of moulds, and states: “Without considering all of the relevant parameters it is not possible to make general conclusions”. That is correct, but unfortunately, under the pressure of economics, including time, new strains will have been developed long before the measurement of all of the relevant parameters are complete.

The second reason is a problem of genetics. The genetics of the production of an enzyme in microorganisms, including fungi, is far more developed than

the genetics of morphology where many enzymes, and their corresponding genes, are involved (see the introduction to Sects. 4 and 4.3.1). What is wanted is a kind of Holy Grail: a mould with the morphology of yeast and without loss of inherent production capacity.

The third reason is a screening problem. Mass screening for a better productivity is relatively easy, there are many assays and hundreds of thousands of new, mutated strains can be produced and tested per year for any given product. However, no assays that perform mass screening for a better morphology can handle so many mutant strains.

6 Conclusions and Prospects

1. Powerful new research tools have been developed, resulting in greater insight into the morphological details of fungi and an increase in the quantity of information and its quality.
2. Highly structured models exist for morphogenesis and for the relation between morphology and productivity. The application of these models in industry seems to be rather limited for a number of practical reasons.
3. The mechanistic postulates underlying these models vary from some that are firmly based on sound physiological principles to others that are in conflict with these principles.
4. Whether or not it is important that a model have a sound mechanistic background depends on the purpose of the model and whether or not it will be used for extrapolation.
5. The physiology of the morphogenesis of fungi is making progress, but the know-how of the genetics behind it is very limited. This is most unfortunate, because a sound genetic base is very important for the future development of fungi with a high inherent productivity combined with morphological properties that result in high rates of momentum, mass, and heat transport.
6. Mass screening and enrichment cultures for favourable morphologies may possibly fill the gap until the basic genetics for morphology has been developed sufficiently far.
7. The control of morphology should be based on insight into genetics, physiology and biochemical engineering, and on real integration of those three areas, in particular among the people working in each of these areas on one project.
8. Other methods of overcoming the morphology problems might be:
 - a. Growing fungi on carrier particles in the form of pellets or as layers. This has been the subject of many studies, but here too the application in industry is limited.
 - b. The use of other microorganisms, such as yeasts or bacteria.
 - c. The use of moulds in solid state processes.
 - d. The use of plants [68, 69]. This point needs some explanation. Production of microbial enzymes, such as phytase, in plants has been proven to be feasible. Phytase is a very interesting enzyme for the manufacture of fodder for pigs and poultry, because it reduces the phosphate content of the

manure considerably. The enzyme was originally produced in *Aspergillus*. The expression of the phytase gene in plants can be made tissue-specific. Its expression in seeds results in a product with relatively high enzyme concentrations. The product is very stable, free-flowing and non-dusting. This method has a broad and very interesting potential for applications.

9. To end with a special remark. The school of Trinci has been standing like a beacon in the landscape of morphology of fungi for a number of decades.

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Appendix

To show how little influence the kind of model can have on the outcome of a simulation, two very simple models for the growth of hyphae due to the transport of vesicles to the tip are compared. In one model, transport is based on diffusion; in the other model, transport is based on flow.

Diffusion Model. The assumptions are: transport of vesicles takes place by ordinary diffusion, and the vesicles are formed with 0th order kinetics. The rate of formation is given by Prosser and Trinci [11]: $r_l = 1.5 \text{ vesicles } \mu\text{m}^{-1} \text{ min}^{-1}$. The value of r_l is not important; the important point is that r_l is constant. The hyphal element consists of one hypha. In the middle of the hypha the concentration gradient of the vesicles is zero; at its tip their concentration is zero, due to rapid uptake of vesicles by the wall of the apex.

The assumptions lead to the following differential equation and boundary conditions:

$$-ID \cdot \frac{d^2C}{dx^2} = \frac{4}{\pi \cdot d_h^2} \cdot r_l \quad (7)$$

The boundary conditions are $dC/dx = 0$ for $x = 0$ and $C = 0$ for $x = L$

The solution of this equation is:

$$C = \frac{2 \cdot r_l}{\pi \cdot d_h^2 \cdot ID} \quad (8)$$

Flow Model. The assumptions are: transport of vesicles takes place by constant flow. The rate of formation of vesicles is identical to that in the diffusion model. In the middle of the hypha the concentration is zero.

These assumptions lead to the following differential equation and boundary condition:

$$v \cdot \frac{dC}{dx} = \frac{4}{\pi \cdot d_h^2} \cdot r_l \quad (9)$$

The boundary condition is $C = 0$ for $x = 0$.

The solution is:

$$C = \frac{4 \cdot r_l}{\pi \cdot d_h^2 \cdot v} \tag{10}$$

For both equations, the flow at $x = L$ is equal to $r_l \cdot L$. That is as it should be, because this is the total amount of vesicles formed per second over the length L , and – in the steady state – it should be equal to the amount arriving at $x = L$. Because the rate of growth of the hypha is proportional to $r_l L$, both models predict exactly the same rate of growth. Needless to say, this is independent of the value of ID or v !

These models can also be used to calculate the apparent velocity due to diffusion of the vesicles. If one calculates the diffusion coefficient ID with the Einstein equation and assumes that the observed transport velocity for a diffusion process can be calculated with

$$v_{observed} \cdot C = -ID \frac{dC}{dx} \tag{11}$$

where C and dC/dx can be calculated from Eq. (1), then the observed velocity is of the same order of magnitude as that found by Prosser and Trinci [11], about $10 \mu\text{m min}^{-1}$.

It is clear that we cannot conclude which model is the correct one from a mechanistic point of view by comparing the simulated rates of growth or the transport velocities with the experimental values. However, if we look at the concentration gradients of vesicles, as observed by Collinge and Trinci [70], then the picture changes completely. In fact, the values of $C(x)$ calculated

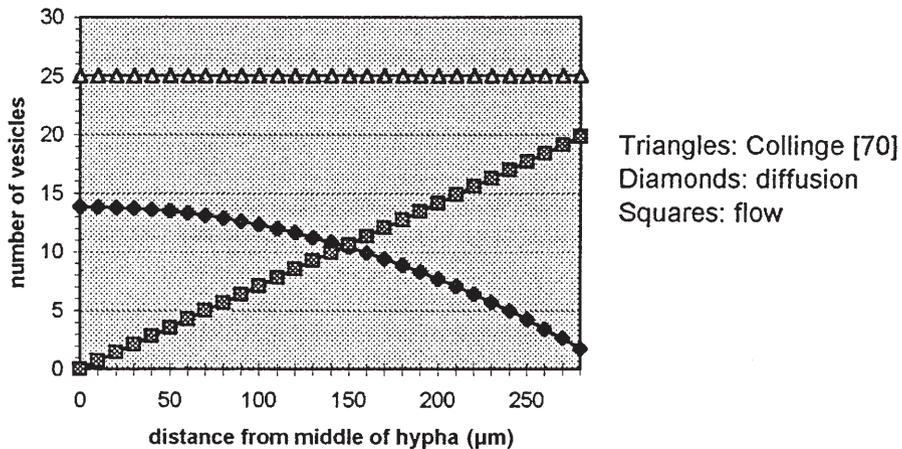


Fig. 7. Concentration of vesicles (number/ μm^3) as a function of the distance to the middle of the hypha in μm .

by both models conflict with the measurements of $C(x)$ for *Neurospora crassa* where $C(x)$ is almost constant at about 25 vesicles per μm^3 until the apical compartment has been reached, where it increases sharply.

Both models presented above are gross oversimplifications of reality and are therefore not very realistic. However, the message is clear: without close observation of reality, the models for the growth of hyphae cannot be considered to be mechanistic in the physical/physiological sense.

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Antibiotica Research in Jena from Penicillin and Nourseothricin to Interferon

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Milestones of antibiotics research and biotechnology in Jena/Thuringia are: 1938 – Hans Knöll established a strain collection of microorganisms; 1942 – production of penicillin on laboratory scale by Hans Knöll; since 1945 – development of industrial production processes for penicillin and streptomycin; 1952 – production of BCG-vaccine; since 1956 – development of biotechnical processes in the Institute of Microbiology and Experimental Therapy for actinomycin C, oxytetracyclin, erythromycin, paromomycin, turimycin, griseofulvin, nystatin, and nourseothricin, and in the 1980s for streptokinase, staphylokinase, and interferons. After the German unification the Hans-Knöll-Institute for Natural Products Research was founded.

Keywords. Bioprocess development, Penicillin, Streptomycin, BCG-vaccine, Nourseothricin, Lysin, Streptokinase, Staphylokinase, Interferons.

In 1937 the well-known glass factory Jenaer Glaswerk Schott & Gen. started cooperation with the young physician Dr. Hans Knöll, living in Frankfurt (Main), in order to check their all-glass bacterial filters. These filters were produced for the first time in 1935 according to an invention of Dr. Paul Prausnitz, Head of the Department for Design and Manufacture of Apparatus, in this glassworks. Knöll had already dealt with the problems of tuberculosis and chemotherapeutics. He subsequently became known for the filtration of bacteria. On behalf of this company, Knöll developed an accurate measuring procedure for checking such filters. Being very interested in this, the Schott-factory offered him the opportunity to establish and manage a bacteriological laboratory in the glassworks. Knöll started this job on 1st November 1938, not being aware at this time that his work would become of special importance both for him and for future biotechnology activities in Jena.

Knöll established a still existing collection of defined strains of different microorganisms as a basis for filter checking, and further works in the fields of microbiology, chemotherapy, and cell biology. His activities in identifying new methods in phase contrast and fluorescence microscopy led to cooperation with the precision-mechanical-optical factory Carl Zeiss in Jena.

Through reference to the literature, Knöll's attention was drawn to penicillin, which had been discovered by Fleming in 1928 and which had been isolated by a research team in Great Britain in 1939. He started experiments to obtain this new antibiotic in mold cultures of *Penicillium notatum*. After a short time crude samples of this antibiotic were obtained. Originally Knöll intended to test penicillin for its effectiveness against cancer cells. However, the immense importance of penicillin in the fight against several human bacterial infectious

diseases soon became known. Penicillin-producing mold strains were sought in the environment and were cultivated in flat glass flasks such as Fernbach flasks. The use of Schott-borosilicatglass proved to be advantageous. It was found that the use of cheaper glass-types spoiled the synthesis of this antibiotic because of its arsenic content. Because of this, and in spite of wartime, an effective exchange – via foreign countries – of information about penicillin took place between Schott-glass and Jena.

Soon afterwards, penicillin wound powder was available from Jena on the laboratory scale. In late 1942, for the first time, it was applied to man. A factory worker, who had a suppurating injury to his hand, was cured successfully by application of this penicillin produced in the Bacteriological Laboratory.

In 1944, this successful work, together with support from the Carl Zeiss factory, led to the transformation of the Bacteriological Laboratory, whose staff had increased from 4 to 15 employees, into the Institute of Microbiology (Schott-Zeiss-Institute), supported by the founding firms of Schott and Zeiss.

At the end of the Second World War in June 1945, the US Army administration at that time in Jena intended to transfer the Institute of Microbiology to the western part of Germany. However, this intention was not realized as in July 1945 the occupation by the Soviet Army began. This new military administration ordered an immediate expansion in penicillin production. Cultivation of the producing mold had been intensified, while stage fermentors made from glass, aluminum, and steel, respectively, were installed in an empty factory building belonging to the firm Carl Zeiss. Railway tank wagons were also modified into fermentors. A great deal of effort was put in to erecting a production plant for penicillin attached to the Institute.

As a result of the rapid increase in the size of the operation, the fermentation section of the Institute of Microbiology was named Jenapharm in 1947. To improve the unsatisfactory supply of medicines, other medicaments and drugs, such as vitamins, analgesics, and transfusion solutions were incorporated into its production program. The number of employees grew to more than 800. Finally, in 1950 the Institute of Microbiology became an independent nationally owned factory, the VEB Jenapharm (VEB means a state-owned company). In addition to penicillin, the VEB Jenapharm produced another antibiotic, streptomycin, which was used to fight tuberculosis, incidence of this illness having increased considerably as a result of war.

A few months after the foundation of the German Democratic Republic (GDR), as a measure in the fight against tuberculosis, the Ministry of Health ordered Knöll to start the immediate production of vaccines according to the methods of Calmette and Guérin as a prerequisite for the introduction of BCG-vaccination. The first research building on the Beutenberg Hill in Jena was therefore erected as a production unit (see Fig. 1), where from 1952 onwards the BCG-vaccine was produced for the whole of the GDR.

In 1953, Knöll left VEB Jenapharm to become director of the newly founded Institute of Microbiology and Experimental Therapy (IMET), which was built in accordance with his ideas, also on the Beutenberg Hill. This institute was taken over by the German Academy of Sciences (later Academy of Sciences of



Fig. 1. Hans Knöll (1913–1988, founder of the Institute of Microbiology and Experimental Therapy on the Beutenberg and the pharmaceutical industry in Jena) left of Werner Eggenrath (Prime Minister of Thuringia) at the topping-out ceremony of the Microbiology Institute under construction in 1951. Knöll was awarded the National Prize of the German Democratic Republic. His Institute became a refuge for the politically displaced. Today the name of Hans Knöll is synonymous with Jena as are the names of Carl Zeiss and Otto Schott

the GDR) in 1956 and was transformed into the Central Institute of Microbiology and Experimental Therapy (ZIMET). Prof. Knöll was director of the ZIMET up to 1976. During this time the personnel at the institute increased to about 1000 workers.

The task of ZIMET was to work on therapeutics, particularly on microbial agents for use in human and veterinary medicine, and later it took over additional technical tasks. The structure of this institute incorporated all the research requirements necessary under the same roof in order to reach a high level of self-sufficiency. ZIMET was composed of the divisions Antibiotic Research, Biotechnology, Experimental Therapy, Medical Microbiology, Methods and Theory, Molecular Biology and Microbial Genetics, Steroid Research, Environmental Microbiology, and Scientific Engineering.

To maintain a continuous line of investigations from screening and increasing efficiency to testing the isolates and purified final products on animals, it was necessary to install qualified microbial and chemical laboratories as well as an efficient experimental breeding system, technical media preparing groups, and different workshops.

Antibiotics research in all its complexity and relevant applications became an essential task of this institute. In the 1950s further improvement in the production of penicillin and streptomycin, in cooperation with VEB Jenapharm, was its main objective. Later on the research potential was systematically developed, including corresponding fields of basic research.

For many years, both closely cooperating divisions "Antibiotic Research" and "Biotechnology" were mainly involved in the elaboration of specific bioprocess methods and down-stream processing by contract with the pharmaceutical industry of the GDR. Further shared research works were the search for producing microbial strains, and the developing of technical instructions for the biosynthetic production and chemical isolation of various antibiotics and other substances for therapeutic and technical purposes, respectively.

The behavior of microbial production strains in shaking flasks and laboratory fermentors was investigated to optimize process conditions and the composition of media on the basis of process kinetic analyses, as well as to elucidate the importance of certain medium compounds for special types of biosynthesis. For such investigations a new biometric screening method, based on the $(2n+1)$ -spectrum, was developed and used with high efficiency.

In the 1960s basic investigations into growth and product-formation kinetics, as well as metabolic regulation in microorganisms, were started, aiming at scientific progress in the optimization of production methods obtained from models. On model systems, in a number of instances, the bistability of specific product formations of microbial processes was demonstrated successfully. Methods for optimal control of biotechnological processes were developed on the basis of mathematical descriptions of growth, metabolism, and product formation in microorganisms as well as by computer simulation of kinetic models.

In a pilot plant with reactors up to 3 m³ net capacity the biotechnical methods were further adapted to the conditions of particular industrial production. Subsequently, the biosynthetically formed substances were obtained by selected known down-stream processes. Moreover, large numbers of small quantities of microbial agents for experimental purposes were produced there.

A widely recognized way to rationalize the multistage procedures of screening and selection, respectively, was opened by introducing six types of selection machine developed by an automation team of the ZIMET. This so-called Autoselect System includes machines to deliver small quantities of agar, to inoculate colonies, to dilute samples, to punch test plates, to pour sample solutions into the punched holes, and to measure the diameters of inhibition zones on the test plates by an optoelectronic method. Thus, by means of the Autoselect Systems, the number of colonies and samples tested could be considerably increased, and the accuracy of working steps and measurements was remarkably improved. The data were evaluated by computer. The optoelectronic measuring device was adopted by the production program of the factory VEB Carl Zeiss Jena.

The search for new antibiotics producing microorganisms had been a permanent task of the ZIMET, particularly of the Antibiotic Research division. Up to the 1980s a collection with more than 20,000 taxonomically identified, freeze-dried strains with defined antibiotic activity had been established.

New production methods were developed and transferred to industrial production for antibacterial antibiotics, such as actinomycin C, streptomycin, oxytetracyclin, erythromycin, paromomycin, and turimycin, as well as for the antifungal antibiotics griseofulvin and nystatin and the nutritive antibiotic nourseothricin, which was required in animal nutrition. Remarkable results, which were used in the industrial production of nourseothricin, were obtained from basic investigations on the regulating effect of phosphate on secondary metabolism. It was detected, that a regulated feeding with phosphate, liquefied starch, and ammonia, while realizing a phosphorus limitation and simultaneously sufficient concentrations of the other components in the medium, can increase the biosynthetic production of this antibiotic in stirred and intensively aerated bioreactors up to yields of 50 g/l nourseothricin. Furthermore, an economic mode of precipitating this agent, basing on its adsorption to bentonite, was invented and applied in industrial production.

Taking multivalent advantages of growth media, an example was set in the industrial streptomycin processes. By a specific change in down-stream processing, a proteolytic enzyme complex could be obtained in addition to the main product, streptomycin. This by-product, after concentrating in its aqueous solution, was used on the industrial scale in the VEB Filmkombinat ORWO Wolfen (Saxony-Anhalt) for the de-gelatination of photographic films in order to recover silver residues. Furthermore, the important VEB Lederfabrik Weida (Thuringia) used this product as a softening-enzyme in the leather-tanning process.

A new method of processing was adapted and tested successfully in the contamination-free production of L-lysine yielding an important food supplement in animal nutrition. In order to guarantee the required high oxygen transfer rate, deep-stream pilot reactors (0,45 m³ capacity) were used. In cooperation with the Research Center of Biotechnology Berlin the developed processing was transferred to industrial application. In the VEB Gärungschemie Dessau (Saxon-Anhalt) non-contaminated production was realized constantly with yields of 80–90 g/l in a production time of 70 h, using a technical deep-stream bioreactor-device with a capacity of 8 m³, fitted out with a continuously acting sterilization-unit for nutrient media. Such a high yield of L-lysine from a bioprocess without contamination was outstanding at that time.

In the early 1980s the ZIMET was concentrating its efforts on the further extension of biotechnology, particularly in the important field of the development of microbial agents from genetically engineered microorganisms. In this connection, genetic engineering protein technology and process development for rDNA products has been promoted.

The genes coding for the plasminogen activators streptokinase and staphylokinase were cloned and sequenced. Expression studies in a variety of hosts led to the construction expression vectors based on staphylokinase signals. Several bacterial strains producing significant amounts of human interferons alpha 1, alpha 2, and gamma have been constructed on the basis of these vectors. In order to maximize the production of interferons in the *E. coli* cells a high cell density bioprocess for the industrial application was developed in 1985. In 1987, after only two years, the first lots (about 10¹⁰ IU) of purified human inter-

Table 1. Institutes of the Academy of Sciences and of the Academy of Agriculture of the former GDR with R&D in Biotechnology

-
- Zentralinstitut für Mikrobiologie und experimentelle Therapie, Jena
 - Zentralinstitut für Molekularbiologie, Berlin-Buch
 - Institut für Biotechnologie, Leipzig
 - Institut für Biochemie der Pflanzen, Halle
 - Zentralinstitut für Genetik und Kulturpflanzenforschung, Gatersleben
 - Zentralinstitut für Ernährung, Potsdam-Rehbrücke
 - Zentrum für wiss. Gerätebau, Mytron, Heiligenstadt
 - Forschungszentrum Dummerdorf bei Rostock
 - Institut für Biotechnologie, Potsdam
 - Institut für Getreideforschung, Hadmersleben
 - Institut für Phytopathologie, Aschersleben
-

Table 2. Universities with scientific and technical activities in biotechnology

-
- Humboldt-Universität, Berlin
 - Karl-Marx-Universität, Leipzig
 - Martin-Luther-Universität, Halle
 - Friedrich-Schiller-Universität, Jena
 - Wilhelm-Pieck-Universität, Rostock
 - Ernst-Moritz-Arndt-Universität, Rostock
 - Technische Universität, Dresden
 - Ingenieur-Hochschule Köthen
 - Technische Hochschule, Leuna-Merseburg
-

feron alpha 1 were supplied to VEB Arzneimittelwerk Dresden for clinical trials phase I.

The Genetic Engineering Pilot Plant building, erected at the end of the 1980s, houses both a process development base for genetic technologies and a manufacturing unit. These provided researchers with capabilities for producing agents in compliance with applicable GMP regulations and for developing new process steps.

The biotechnological activities in the ZIMET were finished in 1991 by the "Abwicklung" of Academy of Sciences of the GDR. In 1992 the Hans-Knöll-Institute for Natural Products Research (HKI) was founded. It is one of five successor institutes of the ZIMET on the Campus Beutenberg in Jena/Thuringia. The other four successor institutes are Institute of Molecular Biotechnology, Institute of Molecular Biology, Institute of Experimental Microbiology, and Institute of Virology.

Biotechnological research and development in the GDR was concentrated in the Academy of Sciences, the Academy of Agriculture and in the Universities. Tables 1 and 2 give an overview of the Research Institutes and Universities of the GDR. According to the "Einigungsvertrag" of the German unification, all the institutes (Table 1) finished their work at the end of 1991.

Development of Bioreaction Engineering

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In addition to summarizing the early investigations in bioreaction engineering, the present short review covers the development of the field in the last 50 years. A brief overview of the progress of the fundamentals is presented in the first part of this article and the key issues of bioreaction engineering are advanced in its second part.

Keywords. Fluid dynamics, Mass and energy balances, Process monitoring and control, Mathematical models, Metabolic engineering, Expert systems.

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List of Symbols and Abbreviations

a	specific interfacial area
$D_L(r)$	axial liquid-dispersion coefficient
$D_r(r)$	radial liquid-dispersion profiles
$d_{Bl}(r)$	bubble-diameter profile
d_s	Sauter bubble diameter
EDS	energy-dissipation spectrum
$k_L a$	volumetric mass-transfer coefficient
M_L	liquid mixing
MAB	monoclonal antibody
MW_{Pr}	molecular weight of product
$MTS(r)$	turbulence macro time scale profile
mm	motionless mixer
Nu	Nusselt Number (heat transfer)
OTR	oxygen-transfer rate
PI	proportional integral control
PID	proportional integral differential control
P/V	specific power input
PS	power spectrum
pH(z)	longitudinal pH profile
$pO_2(z)$	longitudinal dissolved-oxygen profile
pc	pump capacity
RTD_G	gas residence time distribution
R_X	growth rate, calculated from the OTR
SR	shear rate
SS	shear stress
$T(z)$	temperature profile
Tu	turbulence
$Tu(r)$	turbulence-intensity profile
$TDT(r)$	turbulence-dissipation-time profile
t_c	liquid-circulation time
X	cell-mass concentration, calculated from consumed oxygen
$w_L(r)$	liquid-velocity profile
$w_G(r)$	gas-velocity profile
$w_B(r)$	bubble-velocity distribution
ε_G	gas hold-up
η	viscosity, rheology
μ	specific growth-rate
σ_t	surface tension
σ	specific substrate-consumption rate
π	specific product-formation rate

1 Introduction

The first reports on brewing are over 5000 year old [1], but it was not until 1860 that Pasteur recognised that the alcohol was produced by living organisms in a biochemical process [2a, 2b, 2c]. In 1896, E. Buchner isolated the “fermentation” enzyme from the yeast and identified it [3]. After this time, several fermentation processes were investigated and the corresponding microorganisms were identified. Baker’s yeast and fodder yeast became bulk products and were produced in submerged culture. Citric acid was originally produced in surface culture, but – later on – production was carried out in submerged culture as well [4].

However, the technology of fermentation was adapted to biochemical engineering in connection with the large-scale production of penicillin. The Waldhof-type fermenter, which was used for fodder yeast production, was successfully applied to the production of penicillin in submerged operation. Improved strains and bioreactors were developed [5–9] and advanced operation techniques were applied [10a, 10b] to penicillin production.

During the last fifty years, the biotechnology has had many highlights. Between 1950 and 1970 the main topics were the search for new antibiotics and the improvement of their production, as well as the production and biotransformation of steroids.

In order to redress the lack of proteins in developing countries, single cell protein (SCP) projects were carried out between 1970 and 1980. In western countries, yeasts were cultivated on *n*-alkanes, and – after the oil crisis – bacteria on methanol. In eastern countries, yeast was cultivated on gas oil. These projects peaked in the UK with the large-scale production of bacterial protein (Pruteen) by ICI. However, because the SCP could not compete with the inexpensive soy flour as protein fodder supplement, the projects were not economically successful.

In connection with these projects, the development of large-scale bioreactors, air-lift tower reactors in particular, were promoted.

In parallel to the SCP project, the mass cultivation of algae under non-aseptic conditions, a technology suitable for developing countries, was promoted as well. This project failed because of the resistance in developing countries to the acceptance of protein from algae.

The oil crisis between 1975 to 1985 prompted the conversion to fuel additives of renewable energy sources, such as starch, lignocellulose, and hemicellulose from plants, in addition to increased reliance on coalgas fuel. Again, large national projects for the production of ethanol and butanol were undertaken. The highlight of these projects was the production of ethanol from sugar cane in Brazil. This project too failed for economic reasons. The enzymatic decomposition of natural polymers and their conversion into solvents were also investigated in connection with these projects.

Environmental protection, especially biological wastewater treatment, was the domain of civil engineers. However, for the aerobic treatment of industrial wastewater, huge new bioreactors were developed by chemical engineers between 1975 and 1985. At the same time, biochemical engineers developed new reactors for

the anaerobic treatment of heavily loaded waste-water, because the complex interaction of microorganisms in complex mixed cultures required greater knowledge of microbiology and reaction kinetics. Packed-bed- and fluidized-bed bioreactors with immobilized mixed cultures were used for this purpose.

Except for the biological wastewater treatment, the bulk-product projects were unsuccessful, because they could not compete with the low prices of the agricultural products (SCP) and of naphtha (Gasohol). Therefore, the biotechnological projects were later shifted to the development of high value products. Most of these projects were successful and initiated the development of the new industry based on the Life Sciences.

In the 1970s, projects were initiated for the production and biotransformation of secondary metabolites by plant cells (*Catharanthus roseus*, *Atropa belladonna*, *Digitalis lanata*, etc.) in cultures. However, the plant cells quickly lost their ability to form secondary metabolites in cell culture. Only few projects (e.g., shikonin) were successful. In connection with these projects the development of reactors for the cultivation of shear sensitive cells in highly viscous suspensions were promoted. The investigations with plant cells shifted later to plant breeding and the development of transgenic plants

In the 1970s, insect-cell cultivation was initiated for the production of insect virus (*Autographa californica* nuclear polyhedrosis virus), which is supposed to be used as a bioinsecticide of high specificity. However, owing to its high cost, the endeavour was not realised. At present, these insect cells are becoming more widely used, mainly for the expression of high-value heterologous proteins, using recombinant baculoviruses. Insect cells are especially sensitive to shear. In connection with these projects, cell damage by shear stress and turbulence was investigated.

In 1975, Köhler and Milstein succeeded in fusing an antibody producing B-lymphocyte with a permanent myeloma cell, and were able to propagate them in a continuous culture. This success caused high activity in developing hybridoma cells and the production of various monoclonal antibodies (MABs). Because of the high demand for MABs, production was carried out in large aerated bioreactors, which had been developed especially for MAB production

Starting with naturally existing plasmids, plasmid derivatives were developed in the 1970s, and adapted to the specific requirements of genetic engineering. The construction of expression systems for the production of recombinant proteins is realized by a plasmid host system. The necessary expression-plasmids are coded for the protein product, the transcription control of which is often accomplished with inducible promoters. This development led to the start of various activities on the field of genetic engineering. The stabilization of the plasmid-carrying microorganisms had to be solved, as did the suppression of growth of the plasmid free host. The natural folding of the recombinant proteins had to be maintained. In connection with these processes, strategies were developed for the optimal induction of gene expression and for interruption of the process at the right time.

The cultivation of mammalian cells in medicine has a long story, but only the application of genetic engineering to these cells has made it possible to produce large amounts of therapeutically important post-translational modified pro-

teins. For cultures of mammalian cells new techniques were developed: to protect the cells by low shear aeration and stirring; to reduce cost, by avoiding the use of fetal calf serum in the cultivation medium; and to increase the productivity by high cell density by means of cell-immobilization and membrane-perfusion techniques.

At the present a serious competitor is arising in the form of transgenic animals, which produce and secrete these proteins in their milk.

The formation of high value products by genetically modified microorganisms and animal cells requires highly developed process monitoring and control, in order to maintain the quality and human identity of the proteins. Monitoring the process closely allows more information to be obtained, whereupon better mathematical models are developed and better understanding of the process is gained. This is the field of modern bioreaction engineering.

Bioreaction engineering is practised mainly by chemical engineers, because chemical reaction engineering is one of its platforms [11].

The first biochemical engineering courses were organised by chemical engineering departments in MIT (Mateles et al., 1962), Columbia University, University of Illinois, University of Minnesota and University of Wisconsin in the United States, and at the University of Tokyo (Aiba, 1963) in Japan, and the first books on this subject [12–14] were published by chemical engineers and applied microbiologists [15]. After 1980, a large number of books were published on biochemical engineering (e.g., [16–26]). They provide us with a good overview of the state of the art in biochemical engineering.

2 Fundamentals

Transfer across the gas-liquid interface and mixing of the reaction components in gas-liquid chemical reactors influence the chemical reactor performance considerably. The same holds true for submersed bioreactors.

In large reactors, uniform distribution of the substrate is essential for high process performance. Aerobic microorganisms are often used for production; they have to be supplied with oxygen as well. Therefore, the fluid dynamics of the multiphase system and the transfer processes influence microbial growth and product formation. The turbulent forces, which are necessary for high transfer rate and mixing intensity, damage the microorganisms as well.

Several researchers have investigated multiphase reactors with and without microorganisms. Microbial growth and product formation were investigated in batch, fed-batch and continuous reactors, and their dependence on various parameters were described by means of mass and energy balances and kinetic equations. The reaction of the microbes to the physical and chemical variations in their environment can be explained in terms of the physiology of the microbes. Analytical methods were developed for monitoring the key parameters of the process, and the information gained is used for mathematical modelling, control, and optimization of the processes.

It is necessary to investigate the various relationships between particular variables, before the interrelationship between all of them is considered.

2.1

Fluid Dynamics and Transport Processes

In order to evaluate the interrelation between the fluid dynamics and transport processes in bioreactors on the one hand, and the microbial growth and product formation on the other, it is necessary to carry out systematic investigations with various model systems in different reactors. Fluid-dynamic investigations have mainly been performed in the chemical industry and in chemical engineering departments, with the object of designing chemical reactors, but their results are used for the design of biochemical reactors as well.

Between the first and second world wars, several large chemical companies investigated the performance of stirred tank reactors, but the results were kept secret. Only few publications dealt with this topic before and during the second world war [27–29]. In the fifties and the early sixties, several university research groups carried out similar investigations. The key issues were: power consumption, transport phenomena, mixing processes, and reactor modeling. In this period, industrial research groups were especially active, at Merck [30], du Pont de Nemours [31], and Mixing Equipment Co. [31e], all in the United States, where research in this area was also being performed at Columbia University [31c] and the Universities of Minnesota [32], Delaware [33], and Pennsylvania [8]. Similar studies were being carried out in Japan by S. Aiba at Tokyo University [34] and F. Yoshida at Kyoto University [35], in the Netherlands by van Krevelen at Staatsmijnen [36] and Kramers at TU Delft [37], and in the UK by Calderbank, in Edinburgh [38].

Later, the number of research groups dealing with multiphase reactors increased considerably (Table 1). Bubble-column- and airlift-tower loop reactors were investigated by several authors as well (Table 2). As a result, a large num-

Table 1. The leading research groups that have been dealing with fluid dynamics, transfer processes and mixing in stirred-tank reactors in the last thirty years

C. R. Wilke, H. Blanch	University of California Berkeley	USA
D. N. Miller	du Pont de Namours	USA
J. Y. Oldshue	Mixing Equipment Co	USA
F. H. Deindorfer	University of Pennsylvania	USA
M. Moo-Young,	University of Waterloo	Canada
C. W. Robinson	University of Waterloo	Canada
J. Carreau	Ecol. Poy.Techn. Montreal	Canada
A. W. Nienow	University of Birmingham	UK
J. J. Ulbrecht	University of Salford	UK
H. Angelino, J. P Courdec	CNRS, Toulouse	France
H. Roques, M. Roustan	INSA, Toulouse	France
J. C. Carpentier	CNRS Nancy	France
A. Mersmann	University of Munich	Germany
U. Werner, H. Höcker	University of Dortmund	Germany
P. M. Weinspach	University of Dortmund	Germany
H. Brauer	TU Berlin	Germany
M. Zlokarnik, H. J. Henzler	Bayer Co.	Germany
H. Kürten, P. Zehner	BASF Co.	Germany
H. Ullrich	Hoehst Co.	Germany

Table 1 (continued)

K.D. Kiepkke	EKATO Rühr u. Mischtechnik	Germany
F. Liepe	Inst. F. Strömungstechnik, TU Köthen	E. Germany
F. Yoshida	University of Kyoto	Japan
J. Kobayashi	University of Tsukuba	Japan
T. Kono	Takeda Chem. Ind. Co	Japan
J.M. Smith	TU Delft	Netherlands
D. Thoenes	University of Twente	Netherlands
K. van't Riet	University of Wageningen	Netherlands
J. van de Vusse	Koninklijke Shell Co	Netherlands
A. Fiechter	ETH Zurich	Switzerland
V. Linek	Chem. Techn. Inst. Prague	Czechoslovakia
U.E. Viesturs	Latvian Acad. Sci. Riga	Latvia
M. Raja Rao	IIT Bombay	India

Table 2. The leading research groups that have been dealing with bubble column- and airlift-tower-loop reactors in the last thirty years

Y. T. Shah	Pittsburgh University	USA
M.L. Jackson,	University of Idaho	USA
D.N. Miller	du Pont Namours	USA
G.A. Hughmark	Ethyl Co. Baton Rouge	USA
J.R. Fair	Monsanto Co.	USA
M. Moo-Young, Y. Chisti	University of Waterloo	Canada
C.W. Robinson	University of Waterloo	Canada
M.A. Bergougnou	University of Western Ontario	Canada
H. Kölbl	TU Berlin	Germany
H. Hammer	TH Aachen	Germany
H. Langemann, H.J. Warnecke	University of Paderborn	Germany
W.D. Deckwer, A. Schumpe	University of Hannover	Germany
	University of Oldenburg, GBF	Germany
H. Blenke	University of Stuttgart	Germany
U. Onken, P. Weiland, R. Buchholz	University of Dortmund	Germany
K. Schügerl	University of Hannover	Germany
A. Vogelpohl, N. Rübiger	TU Clausthal	Germany
W. Sittig, W.A. Stein, L. Friedel	Hoechst Co	Germany
H. Zehner	BASF Co	Germany
M. Zlokarnik	Bayer Co	Germany
J.F. Davidson	University of Cambridge	UK
J.F. Richardson	Imperial College London	UK
E.L. Smith, N. Greenshields	University Aston, Birmingham	UK
J.S. Gow, J.D. Littlehails	ICI, Billingham	UK
J. Tramper, K. van't Riet	University of Wageningen	Netherlands
J.J. Heijnen	Gist brocades/TU Delft	Netherlands
Y.F. Yoshida	University of Kyoto	Japan
T. Miauchi	University of Tokyo	Japan
Y. Kawase,	Toyo University	Japan
T. Otake,	Osaka University	Japan
Y. Kato, S. Morooka	Kyushu University	Japan
J.B. Joshi, M.M. Sharma	IIT Bombay	India
J.C. Merchuk	Ben Gurion University	Israel
F. Kastanek	Inst. Proc. Fund., Prague	Czechoslovakia

ber of experimental data in laboratory scale are at our disposal, which allow, for example, the prediction of mixing times and oxygen-transfer rates. However, data for large-scale reactors are still scarce. The results of these investigations are summarized in several books [21, 39, 40]. Stirred-tank reactors have recently been modeled with Computational Fluid Dynamics (CFD) [41–45]. Bubble column reactors were modeled with CFD by solving the Navier-Stokes Differential-Equation System [46–51]. These calculations offer greater insight into the fluid dynamics and transfer processes.

2.2

Macroscopic Total Mass, Elemental Mass, Energy and Entropy Balances

Interrelations between the rates of growth, product synthesis, respiration, and substrate consumption have been studied by the macroscopic balance method. Minkevich and Eroshin [52] developed the degree of reduction concept, which considers the number of electrons available for transfer to oxygen combustion.

Erickson [53], Roels [54], Stouthamer [55], and Yamané [56] have further improved this concept. This method was applied on several biological systems (Table 3). The macroscopic balances provide useful relationships for the analysis of growth and product formation. They allow the prediction of the yield coefficients and efficiency factors, e.g. with different electron acceptors.

2.3

Kinetics of Growth and Product Formation

The early investigations of bacterial growth kinetics were reviewed by Hinshelwood [81]. Empirical investigations indicated that the dependence of cell growth on substrate concentration is the same as that of enzyme kinetics, in which Michaelis-Menten kinetics [82] is generally accepted, and which had been extended to competitive and non-competitive inhibitions and complex enzymatic reactions [83].

Table 3. Application of macroscopic balances to various biological systems

Bakers yeast	[54c, 57–63]
<i>Penicillium chrysogenum</i>	[54c, 64, 65]
<i>Candida utilis</i>	[66]
<i>Escherichia coli</i>	[67–69]
<i>Rhodopseudomonas sphaeroides</i>	[70]
Tetracycline by <i>Streptomyces aureofaciens</i>	[71]
Gluconic acid by <i>Aspergillus niger</i>	[72]
Poly- β -hydroxy-butyric acid by <i>Alcaligenes eutrophus</i>	[73]
<i>Klebsiella pneumoniae</i>	[74]
Conversion of D-xylose to 2,3 butanediol by <i>Klebsiella oxytoca</i>	[75]
<i>Enterobacter aerogenes</i>	[76]
<i>Paracoccus denitrificans</i>	[77]
<i>Propionibacterium</i>	[78]
Several microorganisms	[54c, 79, 80]

Monod recommended an analogous relationship for bacterial growth [84], and applied it to several biological systems. The Monod equation was then extended to special cases of bacterial growth, and relationships were developed to cover product formation as well [85–91]. Continuous cultivation of microorganisms became popular. Mass-balance relationships for steady state and substrate limited cultivation (Chemostat) were published [92–102]. These relationships were used for macroscopic material balances in cultures [54c–80].

2.4

Metabolic pathways

A large number of researchers have participated in the discovery of the metabolic pathways of living cells. In the 1930s and 1940s, the glycolysis and the tricarboxylic acid cycle were recognised [103–106]. Overviews of these investigations were presented in the 1950s and 1960s [107, 108]. The present state of the art has been described by Doelle [109] and by Gottschalk [110]. The results of these investigations, and of careful measurements of the concentrations of the main components during the cultivations, allow quantitative analysis of the metabolic fluxes.

2.5

Process Monitoring and Control

2.5.1

pO₂ and pH Measurement

Since its introduction by Clark [111], the membrane-covered dissolved oxygen electrode and its modified versions have been used widely in the practice of biotechnology. The pH-electrodes with glass membrane are based on investigations of MacInnes and Dole [112]. These sodium-glass membranes are still manufactured and sold under the designation CORNING 015, but modern pH glasses contain lithium oxide instead of sodium oxide and have a much wider measuring range [113].

Temperature, dissolved oxygen, and pH are measured in-situ; the other key process variables are monitored either off-line or on-line.

2.5.2

Biosensors

Biosensors are especially suitable for the analysis of complex culture media. They consist of a chemically specific receptor and a transducer, which converts the change of the receptor to a measurable signal. Enzymes, cells, antibodies, etc., are used as receptors. A good review of the history of biosensor development is given in the book of Scheller and Schubert [114]. Enzymes have been used as early as 1956 for diagnostic purposes. The first transducer was a pH sensor combined with phosphatase [115]. The oxygen sensor was first used by Clark and Lyons [116] as the transducer in combination with glucose oxydase

(GOD). Updike and Hicks were the first to immobilize a (GOD)-receptor in a gel. [117]. Enzyme electrodes were also developed by Reitnauer [118]. The first analytical instrument with immobilized enzyme was Model 23 A was put on the market by Yellow Springs Laboratory [119]. Lactate analyzer 640 La Roche was the next commercial instrument [120]. The first enzyme-thermistor was developed by Mosbach [121], and Loewe and Goldfinch [122] developed the first optical sensor. A bacterium was used as receptor instead of enzyme for alcohol analysis by Divies [123]. Cell organelles were used by Guibault for NADH analysis [124], and synzymes by Ho and Rechnitz [125]. Antibodies were introduced by Janata [126] and receptor proteins by Belli and Rechnitz [127] for biosensors. In the last 15 years, the different types of biosensors were being developed [128]. Their application is restricted to laboratory investigations. They are often used in flow injection analysis (FIA) systems as chemically specific detectors [129, 130]. A short analysis time is a prerequisite for process control. Flow-injection analysis, with response times of few minutes, is especially suitable for on-line process monitoring. Flow-injection analysis, developed by Ruzicka and Hansen [131], became popular in the last twenty years in both chemistry [132] and biotechnology [133].

2.5.3

On-line Sampling, Preconditioning and Analysis

The prerequisites of on-line process monitoring are aseptic on-line sampling, sample conditioning, and analysis. The first on-line sampling systems used a steam flushed valve system, consisting of a sampling transfer-tube from the reactor to the analyser, steam supply, a condenser, and four valves for successively sterilizing the transfer tube, withdrawing the sample, and cleaning the transfer tube. Such systems were used for on-line sampling by Leisola et al. [134, 135]. The medium losses, which were considerable, were reduced by miniaturization [136, 137]. Dialysers were the first cell-free sampling systems [138, 139, 140]. Later on, UF membrane filtration was used for sampling and analysis of low molecular-weight analytes, and MF membrane filtration for sampling and analysis of proteins. The first external cross-flow aseptic membrane module that was integrated into a medium recirculation loop [141] was commercialized by B. Braun Melsungen (BIOPEM®); another system [142] was produced by Millipore. The first internal in situ filter for sampling [143] was commercialized by ABC Biotechnologie/Bioverfahrenstechnik GmbH. A coaxial catheter for cell-content sampling was developed by Holst et al. [144], but it was not commercialized. For gas sampling, silicon-membrane modules can be used [145].

Sample conditioning for the analysis of low-molecular-weight components of the medium consists of cell removal, protein removal, dilution or enrichment of the analytes, correction of pH and buffer capacity, removal of toxic components and bubbles, degassing the sample, suppression of cell growth by growth inhibitors, etc. [146].

Modern on-line monitoring systems offer automated sampling, sample conditioning, and analysis [147–149]. Short sampling-, preconditioning-, and analysis times are prerequisites for process control. The internal in situ sampling

system and flow injection analysis with response times of few minutes are especially suitable for on-line process monitoring. On-line gas chromatography (GC) [150, 151, 152] and high performance liquid chromatography (HPLC) [153, 154] are used for process monitoring as well, but their analysis times are several minutes. Mass spectrometry is used for in-line off-gas analysis [155]. Lately, in situ process monitoring with near-infrared Fourier transform (NIR-FT) spectroscopy [156] and 2D-fluorescence spectroscopy [157] became possible. The present state of bioprocess monitoring has been described by Schügerl [130, 158].

2.5.4

Process Control

The classical low-level automatic controls include analog-, on off, sequence-, and feedback control [159, 160].

Low-level controllers are used for the control of flow (PI), gas pressure (PI), temperature (PID), rotational speed of the stirrer (PI), pH-value, dissolved oxygen (PI) and sequence (sterilization, batch and fed-batch process).

Modern control theory was developed between 1950 and 1960 and was applied in biotechnology in the 1970 s. At the same time, advanced computer hardware, especially microcomputers were being developed. Pioneers in computer control were Armiger and Humphrey [161], Bull [162], Hampel [163], Hatch [164], Jefferis [165], Lim [166], Weigand [167], and Zabriskie [168]. The development of computer control is well represented by presentations in the Congresses on Computer Application in Biotechnology [169–173]. The state of the art of control of bioreactor systems has been described by Wang and Stephanopoulos [174], by Lim and Lee [175], and by Bastin and Dochain [176].

2.6

Mathematical Models

The earliest models related growth to the growth-limiting substrate [85, 177–182], and were extended by including inhibition kinetics. (For a review, see Reference [183].)

Later, models of cell population with segregated and structured models were developed. Tsuchiya et al. [184] classified the mathematical models of microbial populations according to Fig. 1. Segregated models consider the heterogeneity of individuals, whereas structured models take the various cell components into account.

Ramkrishna et al. [185, 186] introduced *cybernetic modeling*, which assumes that the cells choose the possible pathways that optimize their proliferation. Shuler et al. [187, 188] developed large-scale computer models for the growth of a single cell. Other structured cell models, developed by Perretti and Bailey [189, 190], take into account the perturbation of the metabolism that occurs as a result of the introduction of recombinant plasmids. The genetically structured models of Lee and Bailey [191] consider plasmid replication in recombinant microorganisms. Other models deal with the proliferation rate influenced by

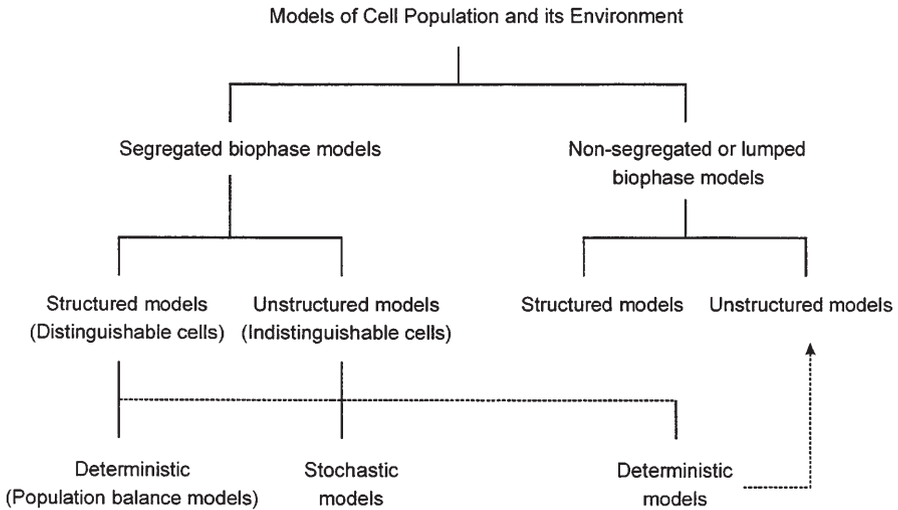


Fig. 1. Classification of mathematical models of microbial population [144]

exogenous growth factors [191]. The metabolic engineering models, recommended by Bailey [192], use the known stoichiometric structure of the intracellular reaction network by assuming a quasi-steady state of the intermediate intracellular metabolites, in order to obtain intracellular fluxes.

However, for process optimization and control, simple structured (so-called compartment) models are used. Harder and Roels compiled common two- and three-compartment models in their review [193].

Several mathematical models, based on the cellular regulation model of Jacob and Monod [194], were developed for the genetic control of enzyme synthesis. These publications were reviewed by Harder and Roels [193]. A typical process model was presented by Bellgardt [195].

3

Interrelation Between Physical, Chemical and Biological Processes

The prerequisites for the determination of the interrelation between physical chemical and biological processes are:

- 1) closely monitored and controlled cultivation;
- 2) monitoring of the key fluid-dynamic properties;
- 3) monitoring of the concentrations of the key medium components;
- 4) monitoring of the concentration and biological state of the cells.

Very few investigations are known that fulfil all of these essentials, but several have been published that satisfy two or three of them.

3.1

Influence of Fluid Dynamics and Transport Processes on Microbial Cultures

Most of the microbial cultivations are performed with monocultures. The important prerequisite is a monoseptic operation, using sterile medium and avoiding infection during the cultivation. The sterility of the bioreactors, necessary for the large-scale production of penicillin, was accomplished by the development of suitable rotating seals for the stirrer shaft. The connections, which are necessary for the fluid-dynamic and mass-transfer measurements, impair the performance of the process and the sterility of the system. Therefore, special setups and runs are necessary for the evaluation of these properties.

The determination of the specific power input (P/V) is only possible for large reactors, because instruments for torque measurement with torsion dynamometer and strain gauges are on the market only for large stirrers. Power monitoring with a wattmeter is only accurate for large reactors, for which the power uptake by frictional losses at the rotating seals of the stirrer shaft are negligible in comparison with the power uptake by mixing and gas dispersion. Therefore, for investigations in laboratory stirred-tank (ST) reactors, power-input data are not available. The specific power input can be calculated from the aeration rate in bubble-column (BC)- and airlift-tower-loop (ATL) reactors. The measurement of the gas hold-up in ST reactors is difficult. In BC and ATL reactors, it can be calculated from the pressure difference between the bottom and the headspace, and by monitoring the liquid level with capacity sensors. Steel and Maxon [196–198] performed the first systematic investigation of the influence of specific power-input on fermentation performance. They investigated the production of Novobiocin by *Streptomyces niveus* in stirred-tank reactors of different capacity (20 l, 250 l, 3000 l, and 6000 l) and with various impellers [196–198]. Their main interest was the dependence of gas hold-up (ϵ_G), oxygen transfer rate (OTR), and productivity (Pr) on the specific power input (P/V), speed (N), and diameter (d_N) of the impeller, the aeration rate (Q_G) and the viscosity of the culture medium (η). In a review, Cooney and Wang compared the OTR and OTR-efficiencies – with regard to power input – of yeast, *Endomyces*, and *Streptomyces* in cultures in different industrial reactors varying in volume from 30 to 128 m³ [199].

Based on the early investigations, the Rushton impeller became the standard stirrer in biotechnology. Only recently, new impellers such as the Scaba agitator and hydrofoil agitators (Lightnin A315 and Prochem Maxflow T), with higher mixing and oxygen transfer efficiencies, have come into use (see [200, 201]).

In the early days of bioprocess technology, bubble column reactors were preferred, because it was easier to maintain their sterility. After the sterile rotation seal for stirrer shaft was developed, ST reactors became the standard reactors for industrial production, because of their flexibility and high performance, especially for highly viscous culture media. AS reactor size increased, ST reactors were replaced by bubble-column (BC)- and airlift-tower-loop (ATL) reactors, mainly in the aerobic wastewater treatment plants of chemical factories. Comparison of these reactor types indicated that ST is a high-performance reactor with low OTR-efficiency with regard to the power input, whereas BC

and ATL are medium performance-reactors with high OTR-efficiency with regard to the power input [202–205]. Therefore, less heat is evolved during the operation in BC and ATL reactors than in ST reactors.

Fiechter and Adler [206, 207] compared the performances of compact stirred-loop (CSL)-reactors with overall volume of 50 l, 550 l, an ATL reactor with 2300 l ATL overall volume, a 100 l torus reactor, and 7 l, 30 l standard STs, by cultivating the yeast *Trichosporon cutaneum*, which is insensitive to glucose repression and does not produce ethanol under oxygen limitation. Therefore, there is a direct relationship between the rates of growth and of oxygen uptake. The growth rate can be calculated from the OTR, and the cell concentration from the consumed oxygen. Xanthan production by *Xantomonas campestris* in a highly viscous medium [204, 208, 209] and *T. cutaneum* cultivation in a medium of low viscosity [210, 211] were used for the comparison of the performances of 30 l, 1200 l ATL/BC bioreactors with 15 l, 300 l, 1500 l, 3000 l ST reactors by Deckwer et al [208–211]. They recommended relationships for the calculation of the volumetric mass transfer coefficient ($k_L a$) as well. Several investigations were carried out with other microorganisms (Table 4). In Table 4

Table 4. Fluid-dynamic investigations with microbial cultivation systems. Stirred tank (ST), Airlift-tower-loop (ATL)-, Bubble-column (BC)-, compact-stirred-loop (CSL)-, and stirred-loop (SL) reactors

Organisms	Reactor	Volume (L)	Measurements	Ref.
<i>Trichosporon cutaneum</i>	ST	7, 30	P/V, OTR, R_X	[206, 207]
<i>T. cutaneum</i>	CSL	50, 550	P/V, OTR, R_X	[206, 207]
<i>T. cutaneum</i>	ATL	2300	P/V, OTR, R_X	[206, 207]
<i>T. cutaneum</i>	Torus	100	P/V, OTR, R_X	[206, 207]
<i>T. cutaneum</i>	BC/ATL	30, 1200	OTR, $k_L a$, R_X	[210]
<i>T. cutaneum</i>	ST	300, 1500, 3000	OTR, $k_L a$, P/V	[211]
<i>Xanthomonas campestris</i>	BC	13	$k_L a$	[203]
<i>X. campestris</i>	ATL/BC	50, 1200	$k_L a$, OTR, MW_{Pr} , η	[204]
<i>X. campestris</i>	ST	1500	$k_L a$, OTR, MW_{Pr} , η	[204]
<i>X. campestris</i>	ST	15, 100	OTR, MW_{Pr} , η	[208]
<i>X. campestris</i>	ST	15, 100	$k_L a$, η , P/V	[209]
<i>X. campestris</i>	SL	900	$k_L a$, η , P/V	[209]
<i>X. campestris</i>	ST	10	OTR, SR, SS	[212]
<i>Escherichia coli</i>	ATL	60	M_L , pO_2 , OTR, $k_L a$,	[205]
<i>E. coli</i>	ST	10	OTR, $k_L a$	[205]
<i>E. coli</i>	ATL	100	$pO_2(z)$, $d_{Bl}(r)$, d , $\epsilon_G(r)$, OTR, $k_L a(r)$, a , $w_L(r)$, $w_{Bl}(r)$, $Tu(r)$, MTS(r), PS, EDS, TDT(r)	[213]
<i>Saccharomyces cerevisiae</i>	ATL	4000	$w_L(r)$, RTD, t_c , D_L	[214]
<i>S. cerevisiae</i>	ATL	250	ϵ_G , t_c	[215]
<i>S. cerevisiae</i>	ATL	4000	RTD	[216]
<i>S. cerevisiae</i>	ATL	4000	RTD $_G$, w_L , D_L , D_G	[217]

Table 4 (continued)

Organisms	Reactor	Volume (L)	Measurements	Ref.
<i>S. cerevisiae</i>	ATL	80, 4000	RTD _G , w _L , ε _G , D _L , D _G , w _B , (in riser and down-comer)	[218 a, b, c]
<i>Gluconobacter oxydans</i>	ST	3.2, 90	M _L	[219]
<i>G. oxydans</i>	ST	25,000	t _c , p _c , k _L a, pO ₂	[220, 221]
<i>Methyphilus methylotrophus</i>	ATL	30	M _L , t _c	[222]
<i>Candida utilis</i>	BC	20	a	[223]
<i>Leuconostoc mesenterius</i>	BC	20	a	[223]
<i>Penicillium chrysogenum</i>	ST	7, 15, 41	M _L , η	[224]
<i>P. chrysogenum</i>	ST	1500	RTD _G	[225]
<i>Aspergillus niger</i>	ST	50	OTR	[226]
<i>Bacillus subtilis</i>	ST	14	pO ₂ (< 100 ppb) butanediol, pO ₂ (> 100 ppb) acetoin	[227]
<i>Neurospora sitophila</i>	ATL	1000	t _c , ε _G , h	[221]
<i>Chaetomium cellulolyticum</i>	BC	40	t _c , ε _G , h	[228]
<i>C. cellulolyticum</i>	ATL	1300	ε _G , η, OTR, k _L a	[229]
<i>N. sitophila</i>	ATL	1300	ε _G , η, OTR, k _L a	[229]
<i>C. cellulolyticum</i>	BC	40	k _L a, η, σ, a (calc.)	[230]
<i>Kluyveromyces fragilis</i>	ATL	120,000	k _L a, OTR, pO ₂ ,	[231]
<i>K. fragilis</i>	ST	15, 100	OTR	[232 a, b, c]
<i>Hansenula polymorpha</i>	ATL	60	pO ₂ (z), , OTR, k _L a, a, d _{BI} , d _S	[232 a, b, c]
<i>H. polymorpha</i>	ATL	60	pO ₂ (z), d _{BI} (r), d _S , ε _G (r), OTR, k _L a, a, w _L (r), w _{BI} (r), Tu(r), MTS(r), PS, EDS, TDT(r)	[213]
<i>Streptomyces niveus</i>	ST	20, 250, 15,000	P/V, OTR, Pr	[196, 197, 198]
<i>S. aureofaciens</i>	ST	112,000	pO ₂ , η, T, k _L a	[233]
<i>S. avermitilis</i>	ST	800	k _L a, OTR, P/V	[200]
<i>S. rimosus</i>	ST	20, 60	Nu, η	[234]
<i>S. albus</i>	ST	20, 60	Nu, η	[234]
<i>Streptomyces</i>	BC/ATL	20,000	M _L , t _c , k _L a, h, P/V, pO ₂ , pH	[235]
<i>Bacillus licheniformis</i>	ST	100, 67,500	k _L a, η, P/V	[236]
<i>Endomyces sp.</i>	ST	20, 40	P/V, η	[237]
<i>Aureobasidium pullulans</i>	ST	8, 14	DOC	[238]
<i>Candida boidinii</i>	ATL	60	ε _G , d _{BI} , d _S , OTR, k _L a, a, P/V, Pr, pO ₂ , (con- current and counter- current operation)	[239]
cell free system	ATL (m.m)	60	w _L , D _L , w _B (small, large horizontal, vertical)	[240]
cell free system	ATL	160	w _L (r), w _G (r), ε _G (r),	[241]

only investigations are listed that were performed with stirred-tank-, bubble-column-, and airlift-tower-loop reactors with volumes of at least 20 l, because the fluid dynamics in smaller reactors considerably differs from those in large ones. In most of these investigations, the temperature, (T), the stirrer speed (N), the aeration rate (Q_G), and the pH-value were controlled, and the dissolved-oxygen concentration (pO_2) was monitored. In addition to the concentrations of the cell mass, the substrate (S), and the product (P), the composition of off-gas (O_2 , CO_2) were measured. Based on the mass balances, these data allow the evaluation of the specific growth rate (μ), the specific substrate-uptake rate (σ), and specific product-formation rate (π), as well as the volumetric mass-transfer coefficient ($k_L a$), the rates of carbon dioxide formation (CPR) and oxygen uptake (OUR) and their ratio – the respiratory quotient (RQ). The yield coefficients of growth and product formation with regard to the substrate consumption ($Y_{X/S}$ and $Y_{P/S}$) are calculated as well. These process variables were determined in several modern fluid-dynamic investigations that were performed during microbial cultivation. The relationships, which were developed on the basis of these investigations, differentiate between low viscous cultivation media (e.g., *T. cutaneum*) [211] and highly viscous non-Newtonian cultivation media (e.g., *Xanthomonas campestris*) [209]. However, this classification only holds true for systems in which the high viscosity is caused by the product (e.g., xanthan), and not by the microorganisms (fungi or streptomycetes). In the latter case, the cell morphology has a considerable influence on the interrelationship between fluid dynamics and transport processes, and – thus – on cultivation performance. Therefore, no generally applicable relationships exist.

With increasing reactor size, cell concentration, and medium viscosity, the distributions of substrate and dissolved oxygen in a reactor becomes less and less uniform. In large ST reactors, three relatively distinct regions with different mixing intensities exist: (a) a well mixed impeller region (micro-mixer); (b) a slightly mixed bulk region (macro-mixer); and (c) a wall region (dead water). The microorganisms circulate through the micro-mixer – with high substrate and dissolved oxygen concentrations, and the macro-mixer – with low substrate and dissolved oxygen concentrations. Depending on the circulation-time distribution, the microorganisms can become substrate and oxygen limited, respectively, in the macro-mixer region. Bajpai and Reuss considered the coupling of mixing and microbial kinetics in different reactors [226]. In ATL reactors, only the riser is aerated, therefore, oxygen limitation can occur in the down-comer, again depending on the circulation-time distribution in the reactor.

To investigate the dynamic behaviour of the cells in periodically varying environmental conditions, several authors cultivated yeast cells in a small reactor varying the dissolved oxygen concentration periodically and circulating the cell suspension through aerated and nitrogen gassed small reactors, respectively, or adding glucose pulse to the reactor and monitoring the concentrations of cell-mass, ethanol, dissolved oxygen, and NAD(P)H-dependent culture fluorescence [242–249]. The measurements in industrial Baker's yeast ATL reactors indicated that the ethanol, which was produced by the yeast in the down-comer, was consumed in the riser, as long as the volume ratio of riser to down-comer

was large enough [250, 251]. However, in large ST reactors with highly viscous media this phenomenon impairs reactor performance considerably [226].

3.2

Process Identification by Advanced Monitoring and Control

The regulation of metabolic flux and intracellular metabolite concentrations is exerted at various levels, which can be roughly divided into two categories: (a) modulation of enzyme specific activity or affinity to substrates; and (b) modulation of enzyme concentration. Typical examples of the first category include cooperative effects, allosteric effects, and covalent modifications, whereas the second includes transcriptional and translational control [252]. Metabolic control analysis (MCA) had already been developed in the 1970 s [253–256], but the practical application of these results was hindered by experimental difficulties. Twenty years later, the experimental techniques attained the necessary level for the application of MCA. In addition to the standard process variables (T, N, Q_G, pH, X, Pr, S, pO₂, O₂, CO₂), several other variables have to be monitored for metabolite control of growth and product formation. Such variables are the concentrations of phosphate, ammonium, amino acids, primary metabolites (acetate, lactate, pyruvate, succinate, ethanol, etc.), the precursors of the secondary metabolites, NAD(P)H, RNA, DNA, proteins, and cell morphology [130, 257]. The heat generated by the microorganisms [258–263] and the intracellular enzyme activities and metabolites [264–269] are monitored as well. In special cases, the intracellular enzyme activities were measured on-line [270–274].

Using advanced measurement and control techniques in well-mixed reactors, highly reproducible data were obtained [275], allowing pathway analysis in microorganisms, e.g., glucose transport [276], oxygen utilization, and the determination of tricarboxylic acid cycle activity [277]. On-line process identification, by means of elemental and macroscopic balancing and advanced data processing, thus became possible [68, 69, 166, 278, 279].

3.3

Metabolic Engineering, Metabolic Flux Analysis

The regulation of metabolic networks is complex, because they involve several enzymes and a great variety of control mechanisms. Quantitative analysis of intracellular fluxes is an important tool for investigating control mechanisms.

Intracellular fluxes can be determined by measuring the rates of change of extracellular metabolite concentrations and using the total mass and carbon balance to calculate them. The determination of intracellular enzyme activities and metabolite concentrations can supplement these measurements. Particular fluxes along some pathways can be measured directly by NMR spectroscopy using ¹H and ¹³C-, or ³¹P-isotopes. However, direct flux measurements with NMR is impaired by insensitivity. The combination of the mass balances and NMR spectroscopy allows the identification of the critical junctions (nodes) in a network that influence the partitioning of the fluxes, and the intracellular

measurements permit the determination of the type of enzymatic modification. In Table 5, several investigations are compiled.

The extension of metabolite balancing with carbon-isotope labeling experiments allows the quantitative determination of the flux of bidirectional reactions in both directions [306]. The simultaneous application of flux balancing, fractional ^{13}C -labeling of proteinogenic amino acids and two-dimensional NMR-spectroscopy, as well as automatic analysis of the spectra, provides a rapid, double checked analysis of the fluxes [306]. The application of the combination of these techniques led to important new results on the

Table 5. Metabolic flux, metabolic pathway investigations (The NMR-measurements were combined with mass balancing)

Organism	Method	Product	Reference
<i>Bacillus subtilis</i>	mass balancing		[280]
<i>B. subtilis</i>	2D [^1H , ^{13}C]-NMR	riboflavin	[281]
<i>Corynebacterium glutamicum</i>	in-vitro- ^{13}C NMR	lysine	[282]
<i>C. glutamicum</i>	mass balancing	lysine	[283]
<i>C. glutamicum</i>	in-vitro- ^{13}C NMR	lysine	[284]
<i>C. glutamicum</i>	mass balancing	lysine	[285]
<i>C. glutamicum</i>	in-vivo- ^1H , ^{13}C NMR	lysine	[286]
<i>C. melassecola</i>	mass balancing	glutamic acid	[287]
<i>Escherichia coli</i>	mass balancing	(acetate)	[288]
<i>E. coli</i>	mass balancing	(acetate)	[289]
<i>Hybridoma</i>	in-vivo- ^{13}C NMR	MAB	[290]
<i>Hybridoma</i>	in-vitro- ^{13}C NMR	MAB	[291]
<i>Hybridoma</i>	mass balancing	MAB	[256, 292]
<i>Penicillium chrysogenum</i>	mass balancing	penicillin G	[293]
<i>P. chrysogenum</i>	mass balancing	penicillin G	[258, 294]
<i>P. chrysogenum</i>	mass balancing	penicillin G	[295]
<i>Saccharomyces cerevisiae</i> ^a	in-vivo- ^{31}P NMR	(ethanol)	[295]
<i>S. cerevisiae</i>	mass balancing intracell. comp (ethanol)		[297]
<i>S. cerevisiae</i>	mass balancing intra cell. comp. (ethanol)		[298]
<i>S. cerevisiae</i>	mass balancing intra cell comp. (ethanol)		[299]
<i>S. cerevisiae</i>	mass balancing	(ethanol)	[300]
<i>S. cerevisiae</i>	mass balancing	(ethanol)	[301]
<i>Spodoptera frugiperda</i>	mass balancing	DNA, RNA, protein	[302]
<i>Streptococcus lactis</i>	mass balancing	lactic acid	[303]
<i>Zymomonas mobilis</i>	mass balancing intracell. enzyme activity ethanol		[304]
<i>Z. mobilis</i>	in-vivo- ^{31}P NMR	ethanol	[305]
<i>Z. mobilis</i>	in-vivo- ^1H , ^{13}C NMR	ethanol	[306]

^a immobilized cells

distribution of metabolic flux in the cells. Recent results on metabolic engineering are reported in a special issue of *Biotechnology Bioengineering* [307].

3.4

Expert Systems, Pattern Recognition

Expert systems form a special class of artificial intelligence (AI) [308–310]. They handle problems by reasoning, drawing on a knowledge-base generated by human experts who are well versed in the problem in question [311]. Expert systems have been used in chemical engineering [312, 313] and biochemical engineering [314–339]. In Table 6, several biotechnological applications of expert systems are compiled that combine mathematical models with additional heuristic knowledge. Biotechnological processes are highly non-linear and – especially – non-stationary. Hence, their dynamic characteristics change with the time and, consequently, the parameters of the model vary during cultivation. A common way of coping with this problem is to use less detailed models and parameter-adaptive techniques to periodically fit these models to the actual process data. Another way of solving this problem is to use a distributive model – consisting of several sub-models – by which the accuracy of the Extended Kalman Filter used for the estimation of the procedure can be improved, and fuzzy reasoning techniques to incorporate additional heuristic knowledge about the process [332]. By using a modular set of artificial neural networks (ANN), it is possible to obtain reliable real-time state estimations and process predictions [333].

Pattern recognition is used to reconstruct characteristics of the system that are only partially known. Patterns in technological processes are defined as a

Table 6. Application of expert systems

Biological system	References
Baker's yeast cultivation	[314–316]
<i>Bacillus subtilis</i> (α -amylase production)	[317, 318]
<i>Escherichia coli</i> cultivation	[319–323]
<i>Escherichia coli</i> (α -interferon production)	[324]
<i>Lactobacillus casei</i> (lactic acid production)	[325, 326]
<i>Zymomonas mobilis</i> cultivation	[327]
<i>Penicillium chrysogenum</i> (penicillin G production)	[328, 329]
Antibioticum production	[330]
<i>Bacillus amyloliquefaciens</i> (α -amylase production)	[331]
Glucoamylase production	[327]
<i>Fusarium graminearum</i> cultivation	[328]
Beer production	[332–334]
Enzyme production	[335]
Mammalian cell cultivation	[336, 337]
Hybridoma cell (monoclonal antibody production)	[338]
CHO-bioreactor development	[324]
α -interferon recovery and purification	[324]
Pilot plant operation in Eli Lilly Co.	[339]

combination of signals transmitted by different sensors. Each single trajectory occupies a row in the matrix. Hence, this results in as many rows as there are sensors, and as many columns in each row as there are discretized time-steps [342–346]. Pattern recognition provides the possibility of distinguishing many different actual states of a culture, making it a suitable tool for expert systems.

4 Particular Systems

The bioreaction-engineering problems that arise in connection with immobilized cells and high-density cell cultures, as well as with animal and plant cells, differ considerably from those of low-density microbe cultures. Therefore, these systems are considered separately in this section.

4.1 Immobilized Micro-organisms

Microorganisms can be immobilized by growing them on solid support (biofilm formation), by adsorption or colonization in porous supports, or by entrapment within gels, such as synthetic polymers, kappa carrageenan, agarose, alginate, chitosan, cellulose acetate, and collagen. Cell immobilization allows high cell density to be obtained, and high dilution rates can be applied in continuous cultures without cell washout. However, internal gradients in the microbial aggregates can reduce process performance and gas evolution can destroy the gel beads. Biofilms have been used for acetic acid production for a long time. Enzymes were first immobilized by Manecke in 1959 [347] and by Katchalski in 1964 [348]. Mosbach reported the entrapment of enzymes and microorganisms in cross-linked polymers in 1966 [349], and Johnson entrapped fungal spores in solid matrices in 1969 [350]. Mosbach et al. used immobilized micro-organisms for steroid conversion in 1970 [351]; Toda and Shoda used immobilized yeast cells for sucrose inversion in 1975 [352]; Kiersten and Bucke immobilized microorganisms in 1977 [353]; and Fukui and co-workers immobilized microbodies for biosynthesis in 1978 [354]. A review of the early work on immobilized enzymes and microorganisms was published by Wada et al. in 1979 [355] and by Fukui and Tanaka in 1982 [356]. Reviews on immobilized enzymes were presented in 1978/79 [357–359].

Subsequently, several publications appeared on immobilization of enzymes and microorganisms. The early publications on immobilized microorganisms are listed in Table 7. The number of research groups has working in this area has increased since the 1980s. Furusaki and Seki reviewed the reports published in 1990/91 on immobilized microorganisms and on animal and plant cells.

The first industrial application of immobilized enzymes was presented by Chibata in 1978 [357]. Shibatani reported on the application of immobilized biocatalysts in Japan in 1996 [395].

A central interest in reaction engineering is intraparticle diffusion of substrates, which has been identified as the rate-limiting process. The earliest investigations of substrate-diffusion in spherical gel-particles, using immobilized

Table 7. Early publications on immobilized microorganisms

Microorganisms	Product	Year	Reference
<i>Acetobacter xylinum</i>	glycerol → dihydroxy acetone	1979	[360]
<i>Acetobacter suboxydans</i>	acetic acid	1977	[361]
<i>Acetobacter cells</i>	acetic acid	1982	[362]
<i>Brevibacterium flavum</i>	L-malic acid	1980/83	[363, 364]
<i>Gluconobacter oxydans</i>	gluconic acid	1981	[365]
<i>Brevibacterium flavum</i>	glutamic acid	1981	[366]
<i>Brevibacterium ammoniagenes</i>	L-malic acid	1976	[367]
<i>Pseudomonas dacunhae</i>	L-alanine	1980	[368]
<i>Brevibacterium ammoniagenes</i>	NADP	1979	[369]
<i>Serratia marcescens</i>	isoleucine	1980	[370]
<i>Achromobacter liquidum</i>	urocanic acid	1974	[371]
<i>Saccharomyces cerevisiae</i>	ethanol	1981	[372]
<i>Saccharomyces cerevisiae</i>	ethanol	1982	[373]
<i>Saccharomyces cerevisiae</i>	ethanol	1980	[374]
<i>Saccharomyces cerevisiae</i>	ethanol	1983	[375]
<i>Saccharomyces cerevisiae</i>	ethanol	1981	[376]
<i>Saccharomyces cerevisiae</i>	ethanol	1981	[377]
<i>Saccharomyces cerevisiae</i>	ethanol	1981	[378]
<i>Zymomonas mobilis</i>	ethanol	1983	[379]
<i>Zymomonas mobilis</i>	ethanol	1981	[380]
<i>Pachysolen tannophilus</i>	D-xylose → ethanol	1982	[381]
<i>Streptomyces catteya</i>	thienamycine	1983	[382]
<i>Escherichia coli</i>	D-tryptophan	1979	[383]
<i>Penicillium chrysogenum</i>	Penicillin G	1984	[384]
<i>Penicillium chrysogenum</i>	Penicillin G	1979	[385]
<i>Bacillus sp.</i>	Bacitracin	1980	[386]
<i>Candida tropicalis</i>	phenol decomp	1979	[387]
<i>Pseudomonas denitrificans</i>	denitrification	1980	[388]
<i>Bacillus subtilis</i>	Proinsulin	1983	[389]
<i>Bacillus subtilis</i>	α-amylase	1978	[390]
<i>Streptomyces fradiae</i>	protease	1981	[391]
<i>Claviceps purpurea</i>	clavine alkaloids	1983	[392]
<i>Claviceps purpurea</i>	clavine alkaloids	1982	[393]

enzymes were published by Kasche et al. in 1971/73 [396, 387]. In the case of immobilized aerobic microorganisms and cells, the rate-determining process is oxygen-transfer into the spherical gel particles. The local oxygen level influences the viability of microorganisms and cells considerably [394, 398–400]. The effect of immobilization on the physiology of the cells has been investigated by several researchers, notably by Doran and Bailey on *Saccharomyces cerevisiae* [401]. Plasmid stability in immobilized microorganisms was investigated as well. Plasmid stability is improved by immobilization [402, 403]. Animal and plant cells were immobilized as well. They are discussed in 4.3.

4.2

High Density Cultures of Microorganisms

There is a general tendency to work with high-density cultures of microorganisms, in order to increase the volumetric productivity of reactors. High density of microorganisms can be obtained by:

- immobilization of microorganisms;
- application of hollow fibre reactors in continuous cultivation;
- reactors with cell recycling or cell membrane retention in continuous cultivation; or
- fed-batch cultivation.

The immobilization is treated in Sect. 4.1. Hollow fibre reactors are mainly used for high-density cultivation of animal cells. The main aim of their application for primary metabolite production by microorganisms is product removal during the cultivation. The hollow-fiber reactors were first applied in 1976/77 [404, 405]. The continuous cell recycle reactor, equipped with a centrifuge, had already been applied in 1960s [406, 407]. Margaritis and Wilke used a rotating-membrane filter in 1978 [408]; Rogers used a microchannel-membrane filter in 1980 [409]; and Nishizawa used a cross-flow-membrane module for ethanol production in 1983 [410]. Chang and Furusaki presented a review on early applications of membrane bioreactors [411].

Volumetric yields of lactic acid produced by *Lactobacilli* with immobilized microorganisms, in hollow-membrane, and in cell-recycling membrane reactors, were compared. These investigations indicate that the highest volumetric productivities were obtained in continuous cultures with immobilized cells and cell recycling [158]. The low productivity in hollow fiber reactors is due to the high pH-gradient in the reactor.

The highest cell concentrations were obtained in fed-batch operation. Bauer et al. had already obtained high cell densities in 1976 [412]. Still higher cell densities were obtained by suitable medium-composition and feeding strategies [413–422]. Korz et al. obtained 148 g DCW l⁻¹ in a standard stirred-tank reactor [422] and Märkl obtained 174 g DCW l⁻¹ in a dialysis reactor [423].

4.3

Animal and Plant Cell Cultures

Animal cell culture was developed in medicine towards the end of the 19th century [424–427]. However, modern industrial cell-culture began in the mid-1950s, with the use of animal cells for the research and development of vaccines. Standard cultivation media and cell cultivation techniques were developed in the period 1950–1965 [428, 429]. Propagation of human diploid cells on titanium disks began in 1969 [430]. van Wezel developed microcarriers for primary cells and cell strains in 1967 [431]. At first, these processes employed primary cells; later, normal cells – maintained as cell strains – were approved; at present, transformed cells, which have infinite life spans, are being used. However, the latter are not applied to the production of vaccines and materials for human

use. For vaccine production, BHK cells [432] and other cells [433] are used. The development of animal-cell cultures is followed in the reports of the ESACT meetings [434–437]. Reviews of the development of large-scale cultivation of animal cells have been published by Spier [438, 439] and by Kelley et al. [440]. Special reviews of the early investigations of nerve and muscle, human hepatoma, lymphoid cells, and vaccine production, as well as of the production of tPA, are presented in Vol. 34, and on the production of MAB by hybridoma cells and of β -interferon in Vol. 37 of *Adv Biochem Eng* [441, 442]. Special reviews on early investigations of plant tissue cultures are presented in Vols. 16 and 18 [443, 444] of the same series.

A list of genetically engineered vaccines is given by Ratafia [445].

Small reactors, low specific power input, and membrane aeration are often applied for animal tissue cultures. Therefore, no large fluid-dynamic stress effects are expected. By application of larger ST, BC and ATL reactors and direct aeration, gas-liquid interfacial effects and at high specific power-input the dynamic pressure of turbulence can impair the behaviour of animal cells, because of their sensitive plasma membrane. The mechanisms of gas and liquid dispersion in turbulent field was investigated by several researcher. The dynamic equilibrium bubble and droplet sizes depend on the eddy size or on the Kolmogorov-length-scale of turbulence, $\eta_k = \left(\frac{\nu}{E}\right)$ where η represents the

viscosity and E is the energy dissipated per unit mass. Bubbles and drops larger than this length are unstable and are dispersed. The same holds true for the viability of the cells. Cells on microcarriers, flocs, pellets, and filamentous mycelia are usually damaged, because they are larger than η_k at high specific power-input. However, suspended animal cells are not damaged in directly aerated systems, because of their small size and the low specific power input, i. e. large η_k [446]. In aerated systems, bursting gas bubbles can destroy the cells [447, 448]. Animal cells are harmed by shear forces in pumps and valves, but are less so in indirectly aerated reactors. The effect of turbulent jet-flow on suspension cultures of plant cells was investigated by MacLoughlin et al. [449]. because of the selection pressure, the sensitivity of animal and plant cells against fluid-dynamic stresses gradually diminishes with time.

Anchorage-dependent animal cells grow on microcarriers. Non-adherent cells were first immobilized by Nilsson and Mosbach in 1979 [450] and by Nilsson et al. in 1983 [451]. Plant cells were first entrapped by Brodelius and Nilsson in 1980 [452] and subsequently by Brodelius and Mosbach [453]. Immobilized microorganisms and animal and plant cells were reviewed by Bierbaum et al. [454]. Hollow-fiber and microcapsule bioreactors are currently used for animal cell immobilization as well [455]. The encapsulation of living cells was first accomplished by Lim and Moss in 1981 [456]. Jarvis and Grdina developed a modified version of encapsulation (Encapel) in 1983 [457]. The successful use of hollow-fiber reactors for the cultivation of mammalian cells was first reported by Knazek et al. in 1972/74 [458, 459]. Since then, several other research groups have used hollow fiber systems for mammalian cells. A review of early investigations with hollow-fiber-immobilized mammalian cells was given by Heath and Belfort [455]. Recent results are published in the reports of ESAC meetings [460].

5 Conclusion

The combination of chemical-reaction engineering, biochemistry, and microbiology provide a good platform for the development of industrial production processes. The stoichiometry, the mass and energy balances, and the kinetics of chemical reactions were adapted to biochemical reactions. The standard techniques of chemical process monitoring and control were extended by highly specific enzymatic analyses and control strategies to batch and fed-batch cultivation processes. Process monitoring and control were adapted to immobilized-cell and high-cell-density cultures, as well as to animal and plant cell cultivations. Mathematical models were developed for the complex interrelation between the physiology of the cells and their physical and chemical environment. The discovery of the metabolic pathways allowed deeper insight into cell regulation, and thus to the development of structured mathematical models. Systematic investigations with advanced techniques under reproducible conditions permitted the validation of these mathematical models. Computer-aided process control and optimization became possible by means of well-established models. Reliable mass balances due to advanced process monitoring, intracellular *in vitro* enzyme activity, and metabolite concentration measurements, as well as *in-vivo*-NMR spectroscopy – in addition to the knowledge of the metabolic pathways – made metabolic flux analysis and metabolic engineering possible. Bioreaction engineering has developed during the last fifty years into bioreaction engineering science.

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A View of the History of Biochemical Engineering

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The authors present a view of biochemical engineering by describing their personal interests and experience over the years involving mostly conversion of lignocellulosics into fuels and chemicals and the associated engineering subjects.

Keywords. Biomass conversion, Biochemical engineering, Fuels, Chemicals, History.

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1 Introduction

Biochemical engineering has grown into a very broad subject field. The scope of this article is limited mostly to technology for conversion of lignocellulosic biomass into fuels and chemicals, and the associated biochemical engineering topics. The content reflects the interests or personal experience of the authors. It offers a limited view of the history of biochemical engineering. History, as always, has to be told from many different viewpoints, to achieve an objective and complete exposition.

The phrase, “biochemical engineering”, first appeared in the late 1940s and early 1950s. That was the time shortly after aerobic submerged culture was

launched as a way of increasing the production capacity of penicillin, used to cure battle wounds of World War II (Shuler and Kargi 1992). Fungal mycelia grow naturally on the surface of moist substrates. When mycelia are submerged in liquid nutrients, an adequate supply of oxygen, often in the form of finely dispersed air bubbles to support aerobic biological activities, has become an important requirement. Gas-liquid interfacial mass transfer of oxygen in reaction vessels has since become a challenge to those trained in chemical engineering. An article by Hixon and Gaden (1950) on oxygen transfer in bioprocesses initiated a wave of activities that has often been credited as the first recognition of “biochemical engineering” as an engineering subject requiring systematic studies to understand its governing principles and for acquisition of skills for good design and performance.

Commercial scale biological processing of biomass materials is an activity as old as human civilization. In more recent years, utilization of lignocellulosic biomass has become an actively pursued subject, because of the concerns of future exhaustion of non-renewable fossil fuels. One of the authors of this article, Raphael Katzen, had his first personal experience in wood hydrolysis 60 years ago in the late 1930s (Katzen and Othmer 1942). Since the energy crisis caused by the oil embargo by the OPEC countries in 1974, there have been expanded efforts in research and development aimed at improved conversion efficiency of lignocellulosics as an alternative material resource. The future payoff from successful utilization of lignocellulosics will be enormous. In fact, the future of human society may depend on it, which might not be so obvious to many business leaders and policy makers today, but it will become increasingly clear in the years to come. Most of the lignocellulosic materials today are considered “wastes” or, at best, low value materials. In order to achieve profitable industrial scale conversion of lignocellulosics into chemicals, materials and fuels, concerted efforts of scientists and engineers from many disciplines, including biochemical engineering, are needed.

2

Early Development of Biochemical Engineering

Following the commercialization of penicillin, a large number of antibiotics were also discovered from extensive screening programs by many pharmaceutical companies worldwide in the 1950s and 1960s. There was a strong demand for better designs of aeration systems and deeper understanding of the process of oxygen transfer in biological systems so that many new drugs could be manufactured efficiently. Oxygen transfer became in those years a popular subject for biochemical engineers to engage in. One of the authors of this article, George T. Tsao, spent his early career pursuing this topic, starting with his own graduate thesis. In those years, one of the most important references in oxygen transfer is the comprehensive review by Professor Robert Finn (1954). Two other frequently cited articles on oxygen transfer in bioreaction systems include the one by Hixon and Gaden mentioned above and another one by Bartholomew, Karow, Sfat and Wilhelm. Both, in fact, appeared in the same Fermentation Symposium in 1950.

In studying oxygen transfer, the first problem at the time, was how one could measure its rate. This problem led to another most frequently cited reference by Cooper, Fernstrom, and Miller (1944), where the rate of oxidation of sulfite ions to sulfate ions was described as a method to reflect the rate of oxygen transfer into gas-liquid contactors. While aerobic processes was flourishing in the biochemical industry, aerobic and anaerobic wastewater treatment were also becoming increasingly an important and widely used processes in the sanitation industry. The fundamentals are the same, whether it is a bioreaction medium or a liquid waste, both requiring adequate dissolved oxygen in some stages to support microbiological activities. There was considerable interaction among biochemical engineers and sanitary engineers. For instance, among the early references important to oxygen transfer is the publication edited by McCabe and Eckenfelder (1955) on "Biological Treatment of Sewage and Industrial Wastes."

While the field of biochemical engineering was growing in its infancy, the field of chemical engineering was maturing in the 1960s. The famous "Bird" book on Transport Phenomena (Bird, Stewart, and Lightfoot 1960) started to place a solid scientific foundation underneath many of the chemical engineering processes and operations. Meanwhile, "Chemical Reaction Engineering" had evolved from its chemical kinetics origin into a full and important branch of chemical engineering. The book on this subject by Levenspiel (1962) helped educate several generations of chemical engineers. Ever since, design and performance of chemical reactors as well as bioreactors have had systematic engineering guidelines and principles.

Often, biotechnologists get excited when they find certain super microorganisms capable of synthesis and accumulation of a valuable metabolite. Soon, they realize that the product cannot be marketed and it has to be purified to meet necessary specifications. Bioseparation, a phrase coined much later in the 1980s, also started to become an important branch of biochemical engineering. The early work involved mostly adopting separation techniques such as solvent extraction and crystallization, well developed in chemical process industries, to purify biochemical products such as antibiotics, organic acids, vitamins, and others. It was later in the 1980s, when chromatography, membrane separation, electrophoresis, super centrifugation, and so on, were needed for purifying many protein and other sensitive biological products, that bioseparation started to become an important engineering factor.

In 1959, a new journal entitled "Biotechnology and Bioengineering" was first published by John Wiley, with Professor Elmer L. Gaden as its founding managing editor. This journal has since become one of the most important publications in biochemical engineering. In those days, the word biotechnology meant simply the technology based on activities of biological and biochemical materials. This word still means the same to many people today. However, there are now also many who interpret this word solely as activities related to genetic modification of living systems.

While aerobic processes had a close association with the first coinage of the phrase "biochemical engineering"; anaerobic bioprocesses actually have a long and important history. Wine, rum and whisky making involving ethanol

fermentation have always been the most important bioprocess, where, as every one knows, a large oxygen supply is not desired. Methane generation by degradation of biomass materials, occurring naturally or man-made, is also one of the most important anaerobic processes. Without the anaerobic digestion in the bodies of rumen animals, the meat and agricultural industries would have been very different from what we have today. Lactic acid bioprocesses for either the product or for making silage are again carried out by anaerobic microbes. Many other anaerobic processes are also involved in the preparation of a variety of indigenous foods in different countries.

Besides lactic acid, one of the processes that did become fully industrialized was the bacterial anaerobic bioproduction of solvents (Underkofler and Hickey 1954). For some years, the anaerobic process was the main source of the industrial solvents including acetone and butanol, until the rise of the new petrochemical industry. This strictly anaerobic process now no longer exists except in a few laboratory studies.

3

Early Development on Conversion of Lignocellulosics

The annual production of biomass is 60 billion tons worldwide. Waste biomass in the United States is one billion tons per year. If it can be converted into chemicals and materials, human population will be able to enjoy material abundance for ages to come. Biomass materials are renewable. Their utilization creates no net gains of greenhouse gases in the atmosphere. The processing methods utilize mild reaction conditions, creating relatively few pollutants. Among biomass, lignocellulosic biomass is currently of relatively little use. For instance, together with annual harvest of about 360 million tons of farm crops such as corn and wheat in the United States, there are co-produced about 400 million tons of lignocellulosics such as cornstalks and wheat straws, often referred to as crop residues. Most of these residues are left in the field for natural degradation, or collected and burned. Lignocellulosics typically contain 70% or more by weight of polysaccharides, including cellulose and hemicellulose. Once they are converted into monosaccharides such as glucose, xylose and others, bioprocessing methods can be applied to convert them into a large number of chemicals and fuels. Hydrolysis of cellulose to produce glucose, however, has not been an easy task. Numerous attempts have been made but economic success, even today, is still limited. Processes for cellulose hydrolysis can be roughly divided into three categories: concentrated acids and “solvents,” dilute acids, and enzymes.

3.1

Concentrated Acids and Solvents

The earliest approach to conversion of the carbohydrate fraction to sugars stems from the more than 100 year old Klason lignin determination, in which hemicellulose and cellulose are gelatinized in 72% sulfuric acid, and after dilution with water, hydrolyzed to yield mixed five-carbon and six-carbon

sugars. The residue from this solubilization and hydrolysis of the carbohydrate fractions leaves a residue identified as lignin (TAPPI 1988). Although this residue could contain other materials such as wood oils and ash, modern chromatographic analysis permits identification of the true lignin content of this residual fraction. Attempts were made in recent years to use this analytical procedure as a pretreatment to gelatinize cellulose and hemicellulose, and then hydrolyze it to obtain a high yield of sugars.

Another concentrated acid process was developed in Germany prior to World War II (Bergius 1933). This process utilized 40% hydrochloric acid for the solubilization stage followed by dilution to complete the hydrolysis. Here, recovery of the large amounts of costly hydrochloric acid is essential. However, during World War II the primary stage of this technology was utilized in Germany to convert wood waste to sugars, followed by neutralization of the acid with sodium hydroxide to yield a mixture of wood sugars and sodium chloride, suitable for use as cattle feed, as a partial replacement or substitute for limited and costly grain feeds (Locke 1945).

3.2

Dilute Acids

Prior to World War II, technology was also developed in Germany (Scholler 1935), utilizing a dilute sulfuric acid percolation process to hydrolyze and extract pentoses and hexoses from wood waste. Several installations were built in Germany prior to and during the war. It is estimated that about 50 Scholler type installations were built in the former Soviet Union during and after World War II. Some of the wood hydrolysates were processed by yeast to produce cell mass in what is called the Waldhof system in Germany. Draft tubes were installed to induce air dispersion into the reaction mixture: to supply dissolved oxygen to growing cells. Draft tube aerators similar in design to the original Waldhof system are now still widely in use in bioreactors and also in aerobic wastewater treatment facilities. When Leningrad of the former Soviet Union (now St. Petersburg of Russia) was under siege for two years by the advancing German army during World War II, hydrolysis of lignocellulosics was used as a source of some digestible carbohydrates. There is a Hydrolysis Institute in that city. Its war-time director earned two Lenin Medals, the highest honor in those days.

One plant was also built at Domat-Ems, Switzerland by Holzverzuckerrungs A.G., employing the dilute sulfuric acid method. A substantial facility was designed, built and operated under direction of Raphael Katzen for the Defense Plant Corporation of the U.S. Government during World War II at Springfield, Oregon for processing 300 tons per day of sawdust from nearby sawmills, yielding 15,000 gallons per day of ethanol, utilizing *Saccharomyces* for the process (Harris 1946). This yield of 50 gallons per ton of wood was approximately 50% of the theoretical yield. The indicated loss of sugars and production of furfural from the pentoses, as well as possible reaction with lignin, resulting in formation of tarry residues which, when mingled with calcium sulfate derived from neutralization of the sulfuric acid, resulted in major scaling and blockage

problems. After full-scale production proved capacity and nameplate production, the plant was shut down as being uneconomic after the war in competition with rapidly developing low-cost synthetic ethanol.

Research on improvement of the dilute sulfuric acid process continued at the Tennessee Valley Authority after World War II (Gilbert 1952). Dilute sulfuric acid hydrolysis of wood and other lignocellulosic materials was also investigated at the Forest Products Laboratory of the U.S. Department of Agriculture, in Madison, Wisconsin. A number of publications (Saeman 1945) from these efforts become important references on the subject in later years. Despite all efforts, ethanol yields from wood with dilute sulfuric acid technology did not exceed 60% of theoretical.

3.3

Enzymes

Again, it was the war-time efforts by researchers at the U.S. Army Quartermaster Research Center in Natick, Massachusetts, that led to the discovery of cellulose hydrolyzing enzymes, commonly known as cellulases. It was told that army uniforms made of cotton were biodegraded quickly in tropical places during World War II. Under the leadership of Elwyn T. Reese and Mary Mandels (1975), cellulases were identified as the cause of degradation of cellulose in cotton fabrics. The culture that was isolated as one of the potent cellulases producers was *Trichoderma viride* which was later re-named *Trichoderma reesei* in honor of Dr. Reese. The Natick center continued to provide leadership in cellulase research for many years.

4

Renewed and Expanded Efforts on Biomass Conversion

After World War II, there was a long period of prosperity of about two decades. Consumption of petroleum products increased quickly. There were relatively little commercial and research interests in alternatives to petroleum. The oil embargo in 1973–74 served as a wake up call, which renewed strong interest in utilization of alternative resources. Renewable biomass and coal were looked upon for possible replacement of fuels and chemicals from petroleum. At the time, George Tsao was on assignment at the U.S. National Science Foundation, on leave from Iowa State University, managing several funding programs as a part of the RANN (Research Applied to National Needs) initiatives. NSF supported work on organizing conferences and workshops to identify research needs. Recognizing the need for biomass research, George Tsao invited Professor Charles R. Wilke of the University of California at Berkeley to conduct a conference on the use of cellulose as a potential alternative resource of fuels and chemicals. It took place in 1974 and the proceedings of that conference were later published as a special volume of *Biotechnology and Bioengineering* (Wilke 1975). That conference served an important function in stimulating renewed interest in the conversion of biomass into fuels and chemicals. In 1978, the first of a series of conferences on *Biotechnology for Fuels and Chemicals*

(first it was named Biotechnology in Energy Production and Conservation, but later revised to cover other chemicals also) was organized by researchers of the Oak Ridge National Laboratory, Oak Ridge, Tennessee, under the leadership of Dr. Charles D. Scott. Later, researchers of the National Renewable Energy Laboratory, Golden, Colorado also joined the effort and this series of conferences has been held annually ever since. In May 1999, two hundred scientists and engineers from many countries attended the 21st Conference held in Fort Collins, Colorado. The conference proceedings were first published as special issues of B&B and later by the journal, Applied Biochemistry and Biotechnology. This series of publications has turned out to be probably the most important information and reference source in this field.

Attempts of applying the concept of first gelatinizing cellulose and then hydrolyzing it to produce degradable sugar and then ethanol have been carried out over the years. In the late 1970s, researchers at Purdue University further investigated that concept by using concentrated sulfuric acid, concentrated hydrochloric acid, together with several other "cellulose solvents" such as Cadoxin (Tsao 1978), resulted in the issue of several patents. The concentrated sulfuric acid method was investigated again at the Tennessee Valley Authority (Farina 1991) and the U.S. National Renewable Energy Laboratory (Wyman 1991). Recent work by ARKENOL in California, USA (Cuzen 1997) and APACE in New South Wales, Australia, have hinged on developing novel, economical methods for separation of sugars from acid, and recovery of substantial amounts of diluted sulfuric acid, evaporated to the required concentration for recycle to the gelatinization stage of the process. Both membrane and ion-exclusion technologies have been tested and developed, toward eventual demonstration and commercialization of the separation techniques. During the process of interacting concentrated sulfuric acid and cellulose, there are likely chemical reactions taking place between the acid and the substrates, which could influence the yield as well as the degradability of the monosaccharides so obtained as well as the acid recovery. This possibility should be carefully investigated before the process can be commercialized. The use of concentrated hydrochloric acid first applied in the World War II era for conversion of cellulose to sugars was again investigated by Battelle-Geneva on a pilot plant basis, particularly of separation of the hydrochloric acid and sugars, as well as re-concentration of the hydrochloric acid for recycle.

The early work on dilute acid hydrolysis was also revisited at Purdue University (Ladisich 1979), in New Zealand (Whitworth 1980) and recently at NREL (Wyman 1992). The use of dilute acids under mild reaction conditions were looked upon by those at Purdue to serve two functions: removal of hemicellulose and pretreatment for cellulose hydrolysis. Under very mild reaction conditions, dilute sulfuric acid removed hemicellulose to form a hydrolysate containing mostly xylose, arabinose and other hemicellulose sugars, without attacking the cellulose fraction in the substrate. After removal of the hydrolysate, cellulose left in the solid residues was then subjected to either dilute acid hydrolysis at a higher temperature or treated with a concentrated sulfuric acid for gelatinization and then hydrolysis. This 2-stage acid Purdue Process generated two sugars streams (Tsao, Ladisich, Voloch and Bienkowski 1982).

Because of the mild reaction conditions, the hemicellulose hydrolysate containing xylose was not contaminated by furfural and other degradation products that would inhibit microbial activities in subsequent bioprocesses. The glucose stream from cellulose was not contaminated by pentoses making its utilization straightforward.

There were also renewed and widespread interests in enzymatic hydrolysis of cellulose using cellulases. Researchers at the Rutgers University conducted a successful culture mutation program (Montenecourt and Eveleigh 1977 1978). A super-productive *Trichoderma reesei* RUT C-30 strain was resulted from it as a mutant of the then best enzyme producer, QM9414, from Natick. This C-30 culture has continued to be one of the best cellulase producers even today. Meanwhile, modern enzymology techniques were extensively applied to investigate the properties and the reaction kinetics of cellulases (Gong 1979) leading to much better understanding of how these enzymes work symbiotically in converting cellulose into glucose.

Developments initiated by the Bio-research Corporation of Japan, a partnership of Gulf Oil and Nippon Mining, resulted in two basic patents, one pertaining to production of the cellulase enzyme (Huff 1976), while the other initiated the principle of simultaneous saccharification and degradation (SSF) (Gauss 1976). This milestone invention overcame the very long saccharification period due to glucose feed back inhibition of the cellulase activities. Major improvement developed by Gulf Oil Chemical Research Group (Emert 1980) at the Shawnee, Kansas Research Laboratory, and later by the same group after transfer to the University of Arkansas, included a continuous 48 hour process for production of cellulase enzymes, as well as a method of recycling active enzyme from the resulting fermented beer by adsorption on fresh feedstock (Emert 1980).

One of the recognized research needs in biomass conversion was the use of 5-carbon sugars derived from hemicellulose for improvement of overall process efficiency. Once glucose is obtained from cellulose hydrolysis, there is no fundamental problem of making good use of it. Xylose derived from hemicellulose is a different matter. Most good glucose-degrading yeast cells cannot metabolize xylose. An important breakthrough made by Dr. C.S. Gong of Purdue University was the conversion of xylose into its isomer, xylulose, using a commercial enzyme, glucose isomerase. Xylulose can then be readily fermented by *Saccharomyces* yeast to ethanol (Gong 1981 and 1984).

The mid-1970s was also the time when gene splicing was made straightforward because of the application of restriction enzymes and other advances in molecular biology. The phrase, Genetic Engineering, was coined at that time. After C.S. Gong's discovery, Nancy Ho was appointed to head a Molecular Genetics Group in the Laboratory of Renewable Resources Engineering (LORRE) of Purdue, with the main objective of splicing isomerase gene into *Saccharomyces* yeast cells. The work did not succeed for almost ten years. Later, the effort was re-directed to replace isomerase gene with two genes: one for a reductase and another one for a dehydrogenase for converting xylose first to xylitol and then xylulose. In addition, a gene coded for the xylulose kinase was also transferred into *Saccharomyces* to build a new metabolic pathway to

convert xylose into ethanol via the pentose phosphate cycle. This effort, as well known today, resulted in success (Ho 1997). This long effort started in the late 1970s and was metabolic engineering in nature long before the phrase, "metabolic engineering," was coined in the early 1990s.

There were several parallel attempts worldwide in searching for microbial cultures capable of degrading xylose to ethanol. The search includes both the use of natural occurring cultures and also genetically engineered ones. Most notable was the work at the University of Florida led by Professor L. O. Ingram, which was made famous by the issuance of the US Patent 5,000,000 (1991). The Ingram group created recombinant *Escherichia coli* and an improved recombinant *Klebsiella oxytoca*, achieving 98% of theoretical conversion of five carbon sugars to ethanol. Work being done at the University of Toronto (Lawford 1998) utilizing *Xymomonas mobilis* also appears promising, along with work by others (Wilkinson 1996). The genetically engineered *Saccharomyces* was successfully tested at the NREL Process Development Unit at Golden, Colorado. Since then, new *Saccharomyces* cultures have been created by Dr. Ho's group, with the above mentioned foreign genes fused into the chromosomes, assuring improved genetic stability.

A challenging problem associated with xylose conversion to ethanol by either natural or genetically modified microbes is really in the rate of fermentation. Often the biomass hydrolysates contain both glucose and xylose and possibly other monomeric components. In the process, glucose will quickly be exhausted but the process may require another day or two to complete xylose conversion. In other words, the xylose-degrading capability will increase ethanol yield from lignocellulosics with a larger sugar basis, but the slow xylose turnover rate may actually decrease the productivity of the fermentation vessel, which is usually expressed in the amount of ethanol produced per unit time per unit volume of the vessel. Future work on xylose conversion to ethanol should include this issue under careful consideration to bring true overall process improvements.

When the efforts on gene splicing for xylose conversion to ethanol were taking place, parallel efforts were also made worldwide for conversion of xylose and other sugars into chemicals other than ethanol. Over the years extensive efforts have been made on production of butanediol, furfural, xylitol, lactic acid, SCP from pentoses. Meanwhile, researchers worldwide also branched out from its early concentration on renewable fuels of ethanol and methane to other chemicals including acetic acid, lactic acid, glycerol, fumaric acid, citric acid, malic acid, succinic acid, aspartic acid, bacterial polysaccharides, acetone, butanol, butyric acid, methyl ethyl ketone, just to mention a few in a partial list. If lignocellulosics-based chemical industries are ever to compete effectively and eventually replace crude oil-based chemical industries, integrated synthetic networks are needed to rival the complexity and sophistication of the current petrochemical synthetic networks. Manufacture of ethanol alone as a single product of a processing plant is unlikely to be economically effective, without crediting the fuel energy value of the residues. By-products and co-products of ethanol should be expected in future large processing and chemical manufacturing enterprises based on lignocellulosics feedstocks. This view seems to

be also held by many researchers in this field. A quick search of the literature in the last 20 years, or a look at some 200 articles presented at the recent 21st Conference on Biotechnology for Fuels and Chemicals will give an unmistakable impression that scientists and engineers worldwide have been and still are pursuing actively many other products from biomass in addition to ethanol.

5 Further Advances in Biochemical Engineering

The required engineering expertise in manufacture of ethanol and bulk chemicals is somewhat different from that in producing high priced health products. For low value products, the required efficiency is important in determining the overall process economics. For pharmaceuticals, assurance of the product purity and safety is of top priority. The price of a new drug can always be properly adjusted to give manufacturers the desired profit. For products like ethanol, process efficiency requirements stimulated a great deal of engineering research and development work over the years. In the late 1960s and early 1970s, there was a worldwide perception of the possible shortage of food to feed the world population. The oil companies placed strong emphasis on conversion of hydrocarbons into single cell proteins. The demand of dissolved oxygen to support cell growth with hydrocarbons as the carbon sources is even stronger than that when carbohydrates are the substrate. The problem of oxygen transfer and cell growth also led to a number of new and low cost designs of fermentation vessels. Airlift reactors with either internal or external circulation loops were tested on a large scale (Schuler and Kargi 1992). At that time, dissolved oxygen analyzers were built and first marketed in the late 1960s. With this instrument, the control of adequate supply of dissolved oxygen became much easier. The above mentioned sulfite oxidation method often led to wrong conclusions when the reaction conditions are not well understood and properly controlled (Danckwerts 1970). With a dissolved oxygen analyzer, not only the DO concentration but also the rates of oxygen input into the reaction mixture as well as uptake by cells can be readily determined with dynamic measuring procedures (Mukerjee 1972, Tsao 1968). The use of oxygen analyzers should be considered a milestone advance in the history of biochemical engineering. The interests in conversion of hydrocarbons into SCP quickly disappeared after the oil embargo in 1974. Soon afterwards, with the advances made in DNA recombinant technology and the energy crisis in the mid-1970s, the emphasis of biochemical engineering changed to work on growth of animal cell cultures and purification of proteins and other modern health products as well as the conversion of biomass into chemical and fuel products.

In addition to the above mentioned SSF (simultaneous saccharification and degradation) process, new methods of simultaneous degradation and product recovery (SFPR) processes were focal points of extensive investigation (Cen 1993). Biological agents almost always suffer from product feedback inhibition. The SSF concept avoids the feedback inhibition of cellulase activities. SFPR or SSFPR will help to reduce the feedback inhibition, for instance, of ethanol on yeast cells. For ethanol, a number of techniques were proposed

and tested, including, for instance, membrane separation of the ethanol product (Matsumuro 1986), evaporation of ethanol from processing beer by gas stripping (Dale 1985, Zhang 1992), pervaporation (Mori 1990), solvent extraction (Bruce 1992), and others. Meanwhile, there has been growing interest, as mentioned above, in producing other chemicals from degradable sugars. For acidic products such as lactic acid intended for preparation of polylactide biodegradable plastics, there was work on simultaneous removal of the acid by adsorption on resin columns coupled with the bioreactors (Zheng 1996). Among the many bioprocesses of commercial interest, past or current, the anaerobic acetone-butanol-ethanol process is the most sensitive to product feed-back inhibition. At only about 8 g/l of butanol, the reaction will be stopped completely. This process has a acid-formation phase followed by a solvent production phase. Acetic acid and butyric acid formed in the first phase are later converted into the solvents. Attempts were made to adsorb the acids with selected resins, resulted in two possibilities. One can, in one case, continue to produce the solvents, or one can change the process to producing organic acids instead (Yang 1994a, b 1995).

In order to improve reactor efficiency, multiple stage, fed batch reaction systems with internal cell recycle has been developed for ethanol production in the industry. For reducing energy cost spent in dehydration of ethanol to produce fuel grade products, the technique of adsorption by corn grits was invented (Ladisch 1984). This technique is currently used in industrial production of about 60% of the fuel ethanol in the United States. For increasing reactor productivity, cell recycle to build up to high cell population density has been investigated. In short, many advances in improved engineering of processing techniques and reactor designs have been resulted from a large amount of research and development efforts in the last decade. This type of true engineering work will continue for years to come to support large scale conversion of lignocellulosics into low value chemicals and fuels where high processing efficiency is of critical importance.

Even though the concept of metabolic engineering was not new, the work in this area took off and expanded greatly in the 1990s. With the tools of DNA recombinant technology and protein engineering well in hand, metabolic engineering promises to create many new opportunities in biomass conversion. The new techniques of DNA chips, microarray, and combinatorial techniques will bring biochemical engineering to yet another high level of sophistication, greatly benefiting the biomass technology.

6 Further Advances in Biomass Conversion

Gelatinization by concentrated sulfuric acid was the first technique of pretreatment of wood before high yields of glucose could be obtained in the subsequent hydrolysis (Cuzens 1997). Since the mid-1970s, several other pretreatment techniques have been investigated with varying degrees of success. The results from enzymatic hydrolysis reported by the Natick group were derived from lignocellulosic substrates after ball milling as a pretreatment. Extensive ball

milling apparently can decrystallize cellulose and making it more accessible to enzymatic hydrolysis. However, the large energy expense in balling milling makes the method impractical. Steam explosion is another pretreatment method being extensively investigated (Schell 1991). The high steam temperature associated with the high pressure degrades large fractions of pentoses from hemicellulose making the pretreatment less desirable. Another explosion method was then invented involving the use of compressed ammonia replacing high pressure, high temperature steam. The low temperature ammonia achieves explosion without the side effect of the high temperature in steam explosion (Holtzapfel 1991). In this process, a high level of ammonia recovery is needed to reduce chemical cost. Yet, another method involving carbon dioxide explosion at low temperature has been investigated (Zheng 1995, 1996). An interesting side benefit observed in carbon dioxide explosion is the sharp decrease in cellulose crystallinity by the pretreatment, in addition to exploding the substrate into fine powders. It has been speculated that carbon dioxide being hydrophobic, unlike steam, is capable of entering the crystal lattice of cellulose and causing disruption.

With pretreatment techniques and bioreactor designs well advanced, product recovery, as well as production of many other chemicals besides ethanol being extensively investigated, the entire field of biomass conversion is on the verge of becoming a real competitor in the chemical manufacturing industry. In more recent years, increasing numbers of chemicals that may have been products of petrochemical processing, are being produced by biomass conversion. One well known example is lactic acid (Du 1998). Fumaric acid is another organic acid for which biomass conversion may become the method of industrial production, replacing petrochemical synthesis (Cao 1996). There is now a very efficient method of production of glycerol by degradation of sugars, which may soon become highly competitive against chemical synthesis of this compound (Gong 1999). Many researchers worldwide are searching for a new bioprocess to produce 1,3-propanediol, a monomer needed in the synthesis of new classes of polyesters. The list of such chemicals is growing and the rate may accelerate before long.

Meanwhile, extensive discussion among experts in the field has identified the high cost of cellulases in the current market as an important roadblock to large scale conversion of lignocellulosics. Cellulase products in the current market are mainly designed for application in textile treatment instead of biomass conversion to chemicals. Textile industry, by nature, can tolerate a higher expense of enzymes than the future ethanol industry.

To search for methods of decreasing the production cost of cellulases, recent work has discovered that solid phase processing (SPF) might be useful (Tsao 1999). If oxygen transfer is the most important engineering problem when the liquid submerged fermentation was first introduced, heat dissipation and also oxygen transfer are the main engineering problems in large scale SPF. The work on improving heat dissipation from and oxygen transfer into porous packed beds of moist solid substrates is just beginning, and much additional systematic work is needed to resolve the difficulties. If SPF can be made to perform well, a new era of bioprocess technology might be born much like the period, from

1960s and on, being the new era of submerged processing mode. SPF is definitely not new because making rice wine, sorghum liquor, and soy sauce in the Far East since ancient times has been done by solid phase processing mode. The old technique of SPF needs new improvements to meet the requirements of the modern society. Indeed, there is a great potential in fulfilling them.

7 Concluding Remarks

As stated above, this article is limited to biochemical engineering and biotechnology applied to the conversion of lignocellulosics into fuels and chemicals. Large and important portions of modern biochemical engineering dealing with health products, bioremediation of toxic wastes, and others are left out for other experts to describe.

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Advances in Enzyme Technology – UK Contributions

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Enzyme technology has been a recognised part of bioprocess engineering since its inception in the 1950s and 1960s. In this article the early history of enzyme technology is discussed and the subsequent developments in enzyme isolation, enzyme modification and process technology are described. These creative developments have put enzyme technology in a position of huge potential to contribute to environmentally compatible and cost effective means of industrial chemical synthesis. Recent developments in protein modification to produce designer enzymes are leading a new wave of enzyme application.

Keywords. Enzyme isolation, Enzyme technology, Enzyme immobilisation, Protein engineering.

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1 Introduction

Some 2500 enzymes have been identified to date [1] and currently around 250 are used commercially in various degrees of purity. However, only 25 enzymes account for 80% of all applications mainly in the processing of starch and for use as domestic and industrial detergent additives for cleaning clothes. The application of enzymes for industrial use is perhaps the widest definition of the term enzyme technology. The term has in the past 40 years been variously replaced by biocatalysis, bioconversion and biotransformation. Confusion has arisen where these latter terms may also be used to describe intact microbial cell catalysed reactions as well as isolated enzyme catalysed conversions. In this article I will focus on the use of enzymes (whether used in an intact cell or isolated) as potential catalysts for single step conversions and chart their development over the past 40 years. This has been a field of intense industrial and academic interest (from around 50 publications per year in the 1950s to a steady 750 per year in the 1990s [2]) and I will therefore necessarily be somewhat selective here.

2 Early History

While enzymes in various forms have been used for many thousands of years for the benefit of mankind, the application of enzyme technology to assist industrial and process chemistry is more recent. In the late nineteenth and early twentieth century there were a few isolated reports, but the first enzyme-based reactions of industrial importance were steroid modifications where the enzyme replaced a series of chemical steps required to introduce a hydroxyl group at a specific position [3]. Achieving such specificity is very difficult by conventional chemistry. The catalytic agent used was *Rhizopus nigricans* and the cells were grown in the presence of the reactant. This simple approach proved effective but widespread application was going to be limited to particular cases. For catalytic use it was recognised that ideally an isolated enzyme was required. Leaving aside the need to isolate the enzyme from the cells in the first place, it was also necessary to stabilize the enzyme (now that it was removed from the protective environment of the cell). One method to achieve this is to attach the enzyme to a solid support. Enzyme immobilization on, or in, a support not only provides a catalyst of sufficient size for ease of separation downstream of the reactor, but also keeps the enzyme structure rigid and therefore confers stability. The drive to achieve an immobilized enzyme began in the 1960s [4]. George Manecke described enzyme resins [5]. Malcolm Lilly [6] and Ephraim Katchalski [7] referred to water-insoluble enzyme derivatives and a fourth pioneer, Klaus Mosbach, described entrapped and matrix bound enzymes [8, 9]. However it was at a meeting in 1971 that the term immobilized enzyme was first used and became the standard nomenclature [10]. The 1971 meeting held at Henniker in New Hampshire (USA) from August 9–13 became the first in a valuable series of Enzyme Engineering meetings held under the

Table 1. Advances in enzyme technology

Process	Advance
Growth process	High cell density growth
Enzyme isolation	Improved enzyme expression New enzyme isolation techniques Enzyme immobilization
Biotransformation	Protein engineering Use of non-aqueous media New reactor designs New reactor operating strategies
Catalyst recycle	Enzyme immobilization
Downstream processing	Integration of reaction with product recovery

auspices of the Engineering Foundation (now United Engineering Foundation). These biannual meetings, which are still running today, have formed the backbone of a strong international community in enzyme technology and we owe much to those early pioneers at that first meeting. The meetings aim to cross disciplinary boundaries and have served as an excellent forum for exchange of ideas. The subsequent years have seen a number of developments in enzyme production, enzyme modification and bioprocess technology. These have led to nine significant areas of advancement based on rDNA technology and process engineering research. These advances are listed in Table 1 and I will describe the development of each of these areas in this article.

3 Enzyme Production

3.1 Introduction

Despite early recognition of the need to isolate enzymes for use as biocatalysts, in a number of cases this is not the most effective catalyst form and several conversion types rely on operation in an intact cell (where the isolated enzyme is unstable or use is made of other endogenous enzyme activities (for example for cofactor recycle and regeneration)). There are three modes of operating an intact cell process: growing cell (where the cells are growing while conversion occurs), resting cell (where the cells are metabolically active but not growing while the conversion occurs) and resuspended resting cell (where the resting cells are resuspended in buffer to avoid the problems of product isolation from medium downstream from the reactor). The options are illustrated in Figure 1. Alternatively the enzyme may be isolated from the host organism prior to biotransformation, in order to reuse catalyst efficiently and/or reduce contaminating activities. This is particularly critical where enzymes are used as

Figure 1. Typical flowsheets for the three modes of intact cell-based biocatalysis

Growing cell	Resting cell	Resuspended resting cell
Cell production/ Biotransformation	Cell production	Cell production
↓	↓	↓
Cell recovery	Biotransformation	Cell recovery
↓	↓	↓
Product concentration	Cell recovery	Resuspension
↓	↓	↓
Product isolation	Product concentration	Biotransformation
	↓	↓
	Product isolation	Cell recovery
		↓
		Product concentration
		↓
		Product isolation

catalysts to assist in the synthesis of pharmaceuticals and final product purity will determine application. Regardless of the final biocatalyst form, the process begins with the production of the cell mass.

3.2

Cell production

A key development and significant reduction in the cost of biocatalyst production has come through the application of so called high cell density growth [11, 12] where bacteria can now be grown at large scale up to 100 g dry weight per litre. This is an order of magnitude improvement on previous biocatalyst production methods. Clearly it is necessary to control the growth rate of such processes to ensure that the oxygen demand is not too high, particularly with fast growing cells such as the commonly used strains of *Escherichia coli*. Routinely this is achieved through feeding the carbon source at a predefined rate to prevent oxygen demand outstripping supply. At large scale this has been found to be particularly important. Using such techniques, far higher titres of catalytic protein can be obtained from the cells. For those systems with well understood genetics a choice of host organism has also become possible on the basis of good expression or an operationally robust strain. Additionally the application of rDNA technology has now led to enzymes being induced more easily and cheaply (at large scale) and expressed at far higher levels in the cytoplasm, periplasmic space or even extracellularly. For instance at University College London we have cloned and overexpressed transketolase (for asymmetric carbon-carbon bond synthesis [13]) as high as 40% of the protein in the cell [14]. At such levels of expression inclusion bodies may be formed in some systems. However even when 15–20% of the protein is expressed as the desired enzyme, subsequent purification may be limited to removal of protease activity alone. Potentially such developments herald the advent of direct immobilisation from unpurified homogenised cells. One commercial process (operated by Glaxo

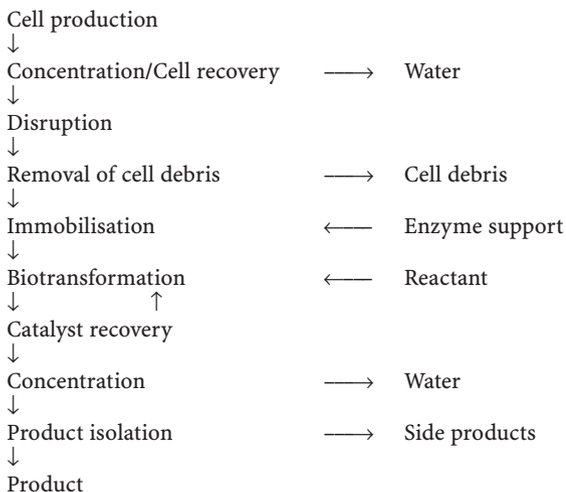
Wellcome in the UK) now uses this for the production of neuraminic acid aldolase for condensation of pyruvate with mannosamine to synthesise *N*-acetyl-D-neuraminic acid (sialic acid) as a precursor to the anti-flu compound, zanamavir [15].

3.2

Enzyme Isolation

In a limited number of cases enzymes are secreted into the growth medium, although at low concentration. However more frequently the useful enzyme activity is intracellular. Figure 2 is a schematic representation of a general flow-sheet for an intracellular enzyme-based process. While a variety of enzyme sources is available, microbial routes are the most productive since the cells are quick to grow, genetics well understood (in some but not all cases), the cells can be broken and the debris separated from catalytic protein effectively. For these reasons most processes commence with a microbial growth process. In those cases where intracellular enzyme isolation is required (for example where transport into the cells is limited or there are competing enzyme-catalysed reactions) the growth process is followed by cell concentration prior to disruption and removal of cellular debris. Isolation of intracellular enzymes was initially done by chemical lysis. This was a difficult process and the ability to disrupt cells via homogenisation [16] enabled cytoplasmic enzymes to be isolated for the first time [17]. This operation has been well characterised and is today standard practice for recovery of intracellular enzyme. The ability to clone into alternative hosts will also have importance here since some cells have been found to be much easier to break open than others [18]. Centrifugation to

Figure 2. Typical flowsheet for isolated intracellular enzyme-based biocatalytic processes



separate the cell debris, prior to purification has also become routine. Entrained air in homogenization and centrifugal operations is a particular problem since globular proteins are very susceptible to shear induced interfacial effects [19]. Surprisingly most enzymes have been found to be resistant to shear itself with the exception of some membrane bound enzymes [20]. Subsequent purification of the enzyme is required only to the extent of removing protease activity or contaminating activities that may affect the final yield of product. This is in sharp contrast to other biotechnological processes where protein purification is a dominant process issue. These techniques now mean that many enzymes can be recovered and isolated effectively for subsequent use as a suspended catalyst or immobilized onto a solid support.

4 Enzyme Immobilization

For intracellular isolated enzyme based catalysis the cost and difficulty of enzyme isolation means that the catalyst becomes a very significant part of the total cost. Hence there is a need to reuse and retain such enzymes. While this was the original justification for enzyme immobilization (necessary also with continuous reactors such as packed or fluidised beds [21]) this is only possible if the enzyme has sufficient stability. Fortunately in many cases the immobilization itself holds the structure of the protein in place and hence leads to significant improvements in the stability of the enzyme. In the 1960s the first work on 'insoluble enzymes' showed that using charged support materials the pH optimum and/or Michaelis constant could be shifted [22] for a particular enzyme. However these electrostatic effects, while academically interesting, did not lead to industrial development due to the high substrate concentrations required for commercial process operation. In 1969 the NSF in the USA began funding of enzyme engineering leading to a considerable interest in the development of immobilized systems. Table 2 lists some of the features of immobilised biocatalysts. One of the earliest applications was by the Tate and Lyle Sugar Company (in the UK) who operated a 6 metre deep bone char im-

Table 2. Features of immobilized biocatalysts

Process operation	Feature
Catalyst production	Loss of activity upon immobilization Additional cost of support
Reactor operation	Reduced activity per unit volume Diffusional limitations Increased stability Protection from interfacial damage ^a Options for alternative reactors ^b
Downstream	Improved separation

^a Gas-liquid and liquid-liquid.

^b Packed and fluidised beds.

mobilized invertase system in the 1940s. The academic research in the 1960s led to higher value processes such as the resolution of racemic mixtures of amino acids by Tanabe Seiyaku (in Japan) [23]. By 1978 at least seven different processes using immobilized biocatalysts were reported to be in commercial operation [24]. The biggest application, and that remains the case today, was for glucose isomerisation (GI) [25]. Since these enzymes are intracellular, justification of the isolation costs was only possible by enzyme multiple recycle. This was achieved via immobilization and used in a packed bed column with downward flow (to minimize residence time and avoid byproduct formation). Previous work had shown the kinetic benefits of operation in packed beds [26] and given very little pH change for the GI reaction an effective process was established. Today the process produces around 8 million tonnes per annum of high fructose glucose syrup. Another key reaction, which is one of the highest value immobilized enzyme processes, is for the cleavage of penicillin G (or V) using penicillin acylase to produce 6-aminopenicillanic acid (6-APA) as an intermediate in the production of semi-synthetic penicillins. Work between University College London and SmithKline Beecham (then Beecham) in the UK developed a process using isolated immobilized enzyme in a substrate fed batch reactor [27, 28]. This enabled pH adjustment to be carried out during the reaction to neutralize the formation of the byproduct, phenylacetic acid, and minimise the severe inhibition of enzyme activity by both products. Substrate inhibition was overcome by feeding. The process was introduced in 1973 and today around 7500 tonnes per annum of 6-APA are made using immobilized penicillin acylase [29]. The process has been much improved since its inception and now operates at an efficiency of about 2000 kg product/kg catalyst. Today many processes use immobilized enzymes and in a limited number of cases immobilized intact cells. In the last decade the use of cross linked enzyme crystals has been examined as an alternative to immobilization [30].

5 Bioprocess Technology

5.1 Introduction

While the power of enzyme catalysis for synthetic process chemistry was recognised in the 1950s, in many cases the properties of a specific biocatalyst were frequently far from ideal for industrial or process use. Since the late 1960s a number of innovative approaches have been applied to processes to overcome these features. Immobilization has been used to create water-insoluble enzyme particles for easy recovery and reuse, operation in organic solvent milieu has led to the effective conversion of poorly water-soluble compounds, reactor choice and judicious operating strategies have minimised reactant and product inhibition, and *in-situ* product removal has minimised product inhibition and toxicity and enabled thermodynamically unfavourable reactions to be carried out. These developments have been crucial to the area and enabled many reactions to be carried out commercially that would not have been possible other-

wise. The techniques are relatively generic and in some cases rules have now been built to assist implementation [31].

5.2

Reactor Choice

Early steroid conversions were carried out with growing cells but it was soon recognised that for many applications the separation of the growth and conversion steps may be beneficial (Figure 1 second column)). Each individual part of the process may be optimised and product recovery effected from cleaner solutions without the presence of complex broth. Operation of a bioreactor and separate biocatalytic reactor also gives the opportunity for resuspension at different catalyst concentrations and use of alternative reactor designs. In other reactions operation with an isolated enzyme is preferable and here there are also choices concerning the optimal reactor configuration. In 1965 Lilly and coworkers [26] published a seminal work on the implications for reactor kinetics on operation in the three classical reactor configurations: batch stirred tank, continuous stirred tank and plug flow reactor. The analysis applied the approach of reaction engineers to evaluate reactors on a kinetic basis (ie which reactor made best use of the available enzyme activity) for enzymes following the Michaelis-Menten kinetic model. The results are presented in Table 3. There are clear advantages for operating in the batch and plug flow reactors with Michaelis-Menten kinetics, where high conversions are required and/or the Michaelis constant is high relative to the initial batch or feed substrate concentration [21, 22]. Another key consideration is the necessity for mixing (required where there is a second liquid phase present, oxygen present, solids present and/or pH control is a necessity) which in many cases will demand operation in a stirred tank configuration. The control of pH, required in any reaction where conversion leads to a change in pH via acid or base consumption or production, necessitates alkali or acid addition to neutralise the effects. Use of buffers is ruled out on the basis of cost, strength (given the concentrations of product required from stoichiometric conversions) and difficulties for

Table 3. Reactor kinetic analysis indicating best (●) use of available enzyme. Choice between batch stirred tank and packed bed is made on the basis of the need for mixing. If Michaelis constant is low and/or conversion required is low there is little difference between reactors

Catalyst kinetics	Reactors		
	Stirred tank		Plug flow
	Batch	Continuous	Packed bed
Michaelis constant high/ High conversion	●	○	●
Substrate inhibition	○	●	○
Product inhibition	●	○	●

downstream product recovery [31]. Consequently most reactors used industrially are batch stirred tanks or fed-batch versions to reduce substrate inhibition. However the introduction of immobilized enzyme as a catalyst form raised the possibility of operation in a plug flow reactor (as a packed bed providing the beads were incompressible). This reactor type not only has kinetic advantages (see Table 3) but also enables a far higher concentration of catalyst within the reactor (voidage around 34% compared to 90% in a stirred tank reactor on account of attrition). For some reactions there has been a clear advantage operating in this way (e.g. fat interesterification and glucose isomerization). In the 1970s a number of reports appeared on the use of fluidised bed reactors [33]. Fluidised beds offer the advantage of high catalyst loading with the possibility of operating with a second phase (liquid, solid or gas). Dependent on the extent of bed expansion, operation can be in plug flow or well mixed regimes. More recently magnetically stabilised beds have been examined as an option to reduce the bead size in a fluidised bed (at the same flowrate) to effect conversions with minimal diffusional limitations [34]. Membrane reactors and a number of other novel configurations have also been examined since the 1980s for multiphasic conversions (and cofactor recycle systems) [35] although commercial application has been limited.

5.3

Medium Choice

While perceived wisdom holds that enzymes operate in water, as early as the nineteenth century there were reports of enzyme catalysed reactions carried out in organic solvents [36,37]. However it was only in the 1970s that the potential of this for enzyme technology was recognised. An isolated report in 1936 [38] was followed by pioneering work in the 1970s by Carrea (in Italy) [39] and Lilly (in the UK) [40]. They argued that the majority of useful compounds for organic chemists are poorly water-soluble and hence, in aqueous-based reaction media, are present at very low concentrations. They introduced water immiscible organic solvents to improve the process by changing the reaction medium [41–43]. Aside from kinetic implications the real benefit lies in higher product concentrations to assist downstream processing. In many cases the product was present in the organic solvent which also assisted isolation. Early experiments involved subjecting cell and enzyme systems to aqueous media which were replaced by organic solvent in increasing proportions.

Klibanov and coworkers [44, 45] took this concept to the extreme by studying the role of water on enzyme catalysis and found that only a fraction of the protein surface need be covered with water for it to remain active. Remarkably this ability to operate in near neat organic solvent led to the possibility of running hydrolytic reactions in reverse (esterification) or even in solvent without water taking part in the reaction (transesterification, interesterification) and a number of processes have now been commercialised. Unilever saw potential in the interesterification reaction for upgrading fats converting mid palm fraction into a substitute cocoa butter and in 1976 described a process using an immobilized enzyme for fat interesterification [46]. The substrate was fed to a

packed bed reactor containing hydrated celite as the support for the lipase. More advanced particles have been introduced subsequently. Maintaining the correct amount of water is critical since too low a level results in a reduced activity and too high a level leads to fat hydrolysis (rather than interesterification) giving a poor quality product containing diglycerides. This reaction exploited the unique properties of lipases which operate effectively at an organic-aqueous liquid-liquid interface. Other enzymes have been shown to work in the bulk of the aqueous phase. However in all cases where organic solvents were introduced the effects were marked, frequently giving increased productivities and better access of the poorly water soluble substrates to the biocatalyst. In the 1980s limitations were also identified for process application. In all but the most insoluble solvents the biocatalyst was denatured to some extent and the amount of water present found to be very critical. An important line of research was commenced examining the role of water on activity [47], the nature of the organic solvent [48,49] and the amount of organic solvent present [50]. Three types of system can be identified: Those in a homogeneous aqueous medium (with water-miscible organic solvent present); those in an homogeneous organic medium (with a small amount of water dissolved in water-immiscible organic solvents) and heterogeneous two-liquid phase systems (water immiscible organic solvent and an aqueous phase forming a separate phase). Each system has found industrial application, although there are particular process reasons for use of a two-liquid phase system [51]. Some representative examples of multiphasic systems are given in Table 4. While lipases function well at an interface and at low water activities, whole cell catalysts require enough water to form a second phase (activity greater than unity). Likewise water may be the substrate or the product of the reaction itself or there may be a shift in relative water solubilities from the substrate to the product. Oxidations are particularly good examples of this where the insertion of the oxygen leads to often dramatic increases in the water solubility of the compound. While in reaction examples such as fat interesterification the use of an organic solvent in the process is a prerequisite for activity, in some cases the addition of an organic solvent may provide an alternative process option. This is well illustrated with a process developed by ICI (now Avecia Life Science Molecules) for the production of *cis*-1,2-dihydro-1,2-dihydroxy-cyclohexa-3,5-diene (DHCD). DHCD may be polymerised to form a precursor to polyphenylene. Polyphenylene has many unique properties and the synthesis of DHCD is chemically intractable. ICI developed

Table 4. Representative examples of multiphasic biocatalytic systems. Where LCA is long chain alkane such as *n*-hexadecane or tetradecane

Aqueous	Medium Organic	Gas	Reaction	Biocatalyst
Water	LCA	Air	Alkene epoxidation	Intact cell
	LCA	Intesterification	Immobilized lipase	
Water	Reactant	Ester hydrolysis	Lipase/esterase	Intact cell
Water	LCA	Air	Aromatic hydroxylation	

a process using a blocked mutant of *Pseudomonas putida* to hydroxylate benzene into DHCD. The reaction was first demonstrated by Gibson and coworkers [52] and the process for biocatalytic conversion first described by Taylor in 1975. While the *cis* glycol product is water soluble the aromatic substrate is only soluble to around 0.5 g/l in water. The substrate can be fed to a batch reactor (although control is difficult) [53] and an alternative is to use a second organic phase. Experiments with addition of tetradecane to form a two-liquid phase system proved particularly successful resulting in up to 60 g/l of product [54]. While the use of organic solvents in biocatalytic reaction media is now finding wide acceptance, debate over the optimal choice of solvent for a particular reaction continues [55, 56]. More recently biocatalysis has been carried out effectively in slurries (with solid substrate and/or product present) and in gas phase media.

5.4

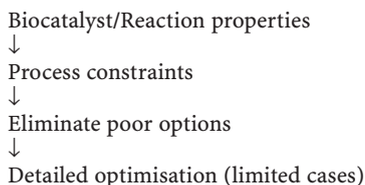
Process Integration

Many biocatalytic reactions are productivity limited by the presence of the product. As the product concentration builds in the reactor it may inhibit the enzyme activity or be toxic to the cell. Likewise reactions which are not thermodynamically favourable may be improved by removal of the product as the reaction proceeds to shift the equilibrium in the synthetic direction. Theoretically such conversions could also be improved by operation with an excess of substrate but this is often precluded on the grounds the substrate is inhibitory to the enzyme, thereby compromising kinetics. *In-situ* product removal (ISPR), where the product is removed from the reactor as it is formed, may therefore provide a tool to overcome some of these problems. Until the 1980s work had focussed in large part on the extractive removal of organic acids and organic solvents from growth processes [57]. More recently extractive ISPR has been examined in relation to the products of enzyme-catalysed reactions [58]. While extraction into an organic solvent gives good capacities for the product being removed, it is not particularly selective. The key need in a biocatalytic reaction is to separate the product from a structurally related substrate. This therefore requires particular selectivity to achieve effective removal [59]. However a number of industrial processes have now used this approach to enable process implementation [60].

5.5

Process Design

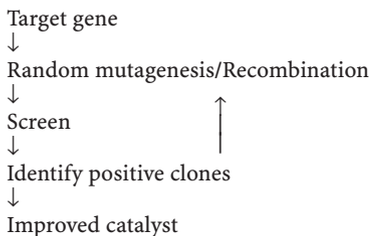
By the mid 1980s it became clear that many enzyme based reactions could be implemented for commercial operation provided one of the growing number of process technology options could be invoked. Woodley and Lilly [31, 61] have pursued the idea of a systematic approach to process choice and selection so as to match the process with the biocatalyst features. An approach was devised based on the simple premise of a relationship between reaction and biocatalyst properties with the final process. This systematic approach (schematically il-

Figure 3. Systematic approach to biocatalytic process selection

illustrated in Figure 3) enables property data to define constraints which then guide process decision making. In essence the approach reduces the number of options for detailed evaluation by eliminating those that are ineffective early on. Part of the rationale for this research was the need to make process design decisions quickly, especially in the pharmaceutical industry where pressure to get to market is great. More recently the representation of the process data has been examined in the form of “process maps” – a graphical technique to visualise the conditions acceptable to the biocatalyst in the reactor [62, 63]. This enables sensitivity analyses and process flexibility along with alterations to the process to be examined using a graphical tool. Combined with high throughput techniques to collect the data to build the map, this will further accelerate the decision making process.

6 Protein Engineering

Most enzymes in nature operate in a highly evolved environment. Unlike chemical catalysts, biological catalysts have evolved over many thousands of years to operate in a narrow range of pH, temperature, ionic strength and concentration. In nature therefore enzymes operate very effectively. However for industrial catalysis three problems are presented. First the enzymes operate at high rates of conversion on natural substrates with the consequence that only low rates are observed with non-natural substrates. This means that conversion of the most interesting, novel, compounds is slow. The second problem concerns the limited range of substrate and product concentrations which give good enzymatic rates and biocatalyst stability. The necessity of efficient downstream product recovery from what is usually an aqueous based medium requires reactor leaving concentrations to be 20–60 g/l for a fine chemical product and an even higher concentrations for lower value industrial (bulk) chemicals. The majority of reactions in nature operate at a mg/l level and hence downstream processing becomes difficult. Thirdly many enzymes operate in a narrow window of pH and temperature. This may not be the same window defined by the substrate and product of a specific reaction. The resulting compromise costs catalytic efficiency and conversion yield. While process techniques can be used to overcome a number of biocatalyst productivity limits, knowledge of enzyme structure (as a result of protein crystallisation developments) in the 1980s

Figure 4. General strategy for directed evolution of enzymic catalysts

raised the intriguing possibility of altering amino acid residues around the active site in order to change the catalyst properties. However the relationship between amino acid changes and the resultant catalytic properties have proved in many cases rather elusive. More recently in the 1990s another approach has been taken. By making alterations in the gene by random mutagenesis (rather than site specific) and/or recombination techniques huge diversity can be generated. This can now be ordered using developments in high throughput screening for favourable properties. Screening these mutants enables selection of positive clones which can be fed back to the mutagenesis stage. In this way the laboratories of Arnold and Stemmer have been able to mimic evolution [64, 65], as illustrated in Figure 4 and preferentially alter biocatalytic properties for industrial exploitation [66]. The work is still in its infancy but the results are impressive, with order of magnitude improvements in specificity, activity and stability in a range of process conditions. The techniques rely on an effective screen and ability to express the genes in a suitable host. Thus far relatively simple proteins have been improved and the goal is to develop appropriate protocols for more complex targets such as redox enzymes [67, 68].

7 Application and Future Perspectives

As discussed earlier the first attempts at commercialisation of enzyme catalysed reactions were in the 1950s and 1960s. It is widely believed that today there are over 100 commercially operating processes from diverse companies. The majority of reactions are in the hydrolytic field primarily with the objective of racemic resolution. This a proven technology for these applications. More interesting however are the academic developments in asymmetric carbon-carbon bond formation and redox catalysis which are now starting to be commercialised and which will change the chemistries to which biocatalysis contributes over the next decade [69–71]. While multistep conversions will see application with intact cell biocatalyst format, for some cofactor-dependent single step conversions the choice between intact cell and isolated enzyme (with cofactor recycle) is becoming a real possibility [72]. The catalyst choice for such conversions will become a particularly interesting area of research in the coming years. Enzyme technology will find a key role in the synthesis of

complex pharmaceuticals and carbohydrates, in particular where optically pure products are required. Their application in the field of large scale bulk chemical synthesis is harder since the economics of these processes means greatly increased product concentrations are required. Nevertheless headway is being made here too and given increasing environmental concerns, the role of biocatalysis is set to increase. While the technology has developed it is clear that there is a need for a greater source of commercialised enzymes. Many new activities are now being identified but there is a need for expression systems in suitable hosts and stable enzymes to be made available to organic chemists. The ability to modify the catalyst and create improved biocatalysts as well as over-express the enzyme, together with effective growth and isolation techniques will be increasingly required by companies in the field.

Enzyme technology has been a field of intense research activity over the past 40 years and it is instructive to note a number of trends in the context of this historical review which point to directions for the future.

- First the range of reactions to which enzyme catalysis can be applied is very broad although currently this is not being exploited. Academic research into carbon-carbon bond formation and redox based reactions will increasingly find industrial application. It is vital that biologists work closely with organic chemists to identify targets.
- Secondly the relative costs which make up the economic analysis of the products of these reactions is changing. Early work was rightly very concerned with yield of product on the catalyst. The catalysts were expensive and not readily available. Since the ability to clone and overexpress enzyme activity, the relative cost of the catalyst has reduced. Alongside this the value of the substrates has increased. Ever more complex substrates with multi-chiral centres are now the starting points for biocatalysis. Hence yield on reactant is becoming increasingly critical. This changes the process analysis and the appropriate technology to enable implementation.
- Thirdly there has been much progress in the implementation of process techniques to overcome non ideal biocatalyst properties. More recently it has become possible to alter the properties of the biocatalyst via genetic and protein engineering. The combination of these approaches will be very powerful and lead to implementation of completely new groups of biocatalysts.

These trends give a perspective on biocatalysis which indicate a bright future providing chemists, biologists and biochemical engineers continue to work effectively together in the way they have come to learn from enzyme technology in the past few decades.

Acknowledgement. Malcolm Lilly, who was to have authored this chapter, sadly died in May 1998. It is therefore dedicated to him as my colleague and friend, whose inspiration and intellectual leadership was in so large a part responsible for the development of enzyme technology.

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Computer Applications in Bioprocessing

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Biotechnologists have stayed at the forefront for practical applications for computing. As hardware and software for computing have evolved, the latest advances have found eager users in the area of bioprocessing. Accomplishments and their significance can be appreciated by tracing the history and the interplay between the computing tools and the problems that have been solved in bioprocessing.

Keywords. Computers, Bioprocessing, Artificial intelligence, Control, Models, Education.

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1 Introduction

To provide some historical perspective about what people were doing with computers and what has changed, I will follow the personalized approach used by others [1]. While pursuing my B. Chem. Eng. and Ph. D. degrees in the late 1940s and early 1950s, I had no contact at all with computers. My thesis was typewritten with carbon copies. After working for more than 7 years at a large pharmaceutical firm where the technical people thought that computers were for payrolls and finance and not of much use for research and development, I joined the faculty of a university in 1963 where about 20% of the engineering professors worked with computers. My education in chemical engineering was not current because my Ph. D. was in biochemistry. I audited a series of five courses in mathematics, studied process dynamics, helped teach it, and thus upgraded my engineering skills.

It was obvious that engineers who used computers could compete better in the real world, so I sought ways to apply computing in both teaching and research. Some professors still rely on their students for any computing, but I felt then and continue to think that you cannot appreciate fully what computers can do when you cannot write programs. I learned FORTRAN but regressed to BASIC when I began to work mostly with small computers. Along the way I have written a few Pascal programs and have dabbled with languages such as Forth. Early in 1997, I switched to Java which presented a very steep learning curve for me because of its object orientation.

I left teaching for another stint in industry from 1973 until 1976. I was in management and ordered a minicomputer for my technical staff. I was the person who used it most but for fairly easy tasks. One program that solved a production problem was for blending of a selection of input lots of stale blood to get adequate values of different blood factors in a product used for standardizing assays in a hospital laboratory. I became fully comfortable with a minicomputer, but my level of sophistication of programming changed little. Our only project related to getting computers into manufacturing tried electronic data logging at the process and carrying the records to the computer for analysis [2].

By the time I returned to teaching, minicomputers were common. The mainframe computer was widely used, but we also had rooms full of smart terminals that were fed their programs from a server. Very soon my research required a computer in the laboratory because we focused on dynamics and control. For

over 20 years we have improved our systems incrementally by upgrading and extending both our hardware and software. All of my graduate students have studied process control, and most have used it in their research. Interfacing a bioreactor to a computer is routine for us, and some of our control algorithms are quite sophisticated. We make some use of artificial intelligence.

2 Historical Development

When I entered academia, analog computers were important. We think of the high-speed of digital computers, but analog computers are lightning fast when handling systems of equations because their components are arranged in parallel. They integrate by charging a capacitor. With large capacitors, voltages change slowly, and the output can be sent to a strip chart recorder or X-Y plotter. Small capacitors give rapid changes with the results displayed on an oscilloscope. Each coefficient is set with a potentiometer, and the knobs can be twisted for testing coefficients while watching the graphs change. This used to be far more convenient than making runs with a digital computer that had essentially no graphical output; the digital results had to be compared as columns of numbers on printed pages. Analog computers have about the same precision as a slide rule, but we are spoiled by the many figures (often insignificant) provided by a digital computer. The Achilles heel of the analog computer is the wiring. Each differential equation requires an integrating circuit; terms in the equation are summed at the integrator's input. Voltages are multiplied by constants by using potentiometers. Constants are developed by taking a fraction of a reference voltage, either plus or minus. With many components, jacks for reference voltages, wires going everywhere for interconnections, jacks for inputs and outputs to pots, jacks for initial conditions, and the like, the hookup for a practical problem resembles a rat's nest. Furthermore, a special unit is needed for each multiplication or division, and function generators handle such things as trig relationships and logarithms. To summarize, analog computers perform summation, integration, and multiplication by a constant very well but are clumsy for multiplication or division of two variables and for functional relationships.

Scaling could sometimes be a chore when setting up an analog computer circuit. The inaccuracy can be great when a constant is not a significant fraction of a reference voltage. Consider, for example, the constant 0.001 to be developed from a reference voltage of 10 V. The pot would have to be turned to almost the end of its range. Proper technique is to scale the constant up at this point and to scale its effect back down at a later point. In addition to magnitude scaling, there can be time scaling when rate coefficients are badly matched.

I spent a fair amount of time with analog computers and enjoyed them very much. I used them for teaching because students could watch graphs change as they tested permutations of coefficients. One terrible frustration with the computers that were used for instruction was bad wires. Students, although admonished not to do so, were thoughtless in yanking wires out of a connection. The wires would come apart inside the plugs where the fault was not

visible. Debugging a huge wiring layout and finding out hours later that one or more of the wires was broken could ruin your day.

I did a little hybrid computing after learning how to do so in a manufacturer's short course. The concept is to let a digital computer control an analog computer. The example most quoted for using a hybrid computer was calculations for a space vehicle. The digital computer was better for calculating the orbit or location and the analog computer, with its parallel and fast interplay, was better for calculating pitch, yaw, and roll. The messy wiring and the difficulty of scaling voltages to match the ranges of the variables doomed both analog and hybrid computation to near extinction soon after digital computers had good graphical output.

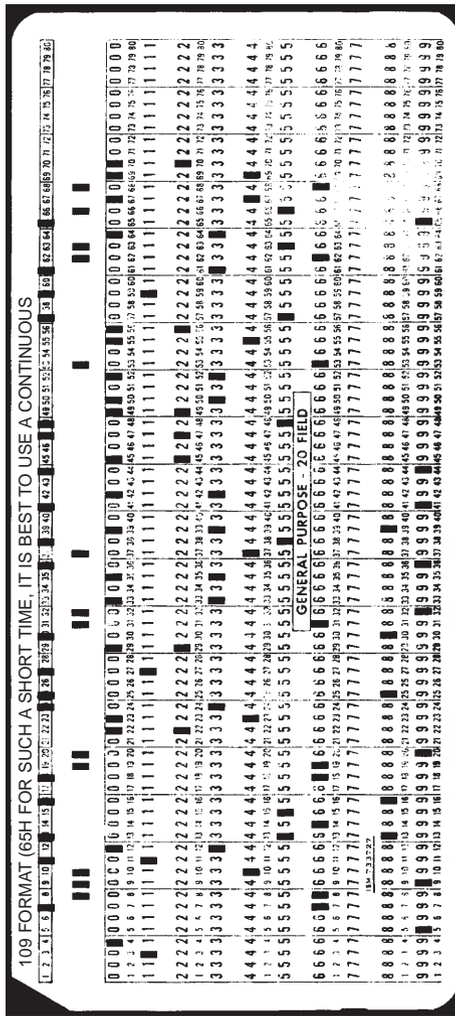


Fig. 1

In the early 1960s, FORTRAN was the most popular language for engineers by far. I learned FORTRAN from books and by examining programs written by others and began to integrate some digital computing into my courses. There were several companies that manufactured main frame computers, and FORTRAN code that I wrote at my university required some modifications before it could execute on another system when I spent the summer of 1970 at a different university.

The IBM punch card was used for communicating with the computer. A typical punched card is shown as Fig. 1. An entire, deep box of cards might be needed to feed the program and the data into the computer. Typical turn around time was overnight, and long runs might not be scheduled for two or three days. Many people were delighted when computer centers could furnish results in an hour or two. Today we have rooms full of personal computers or work stations. In the mid 1960s and through the early 1970s there were rooms full of noisy IBM machines for punching cards. These were fed into a card reader. Wide paper fed on rolls to the printer ended up fan folded with your results. You separated pages along the perforations and held them in thick books with metal strips passed through holes in the paper. There was no graphic output from the printer except when you devised a way to arrange characters as a crude graph. To get real graphs you requested a line printer where a pen moved across the page and touched the paper to make points or lines as the paper was moved back and forth underneath.

Despite the primitive equipment, much could be done. Libraries of code were available for various routine tasks such as a least squares fit of an equation to data points. Remember that the pocket calculator was not common until about 1970 and that mechanical calculators were big, clumsy, noisy, and not very powerful. Feeding punch cards to a computer seemed the best way to calculate even when answers were not ready for a few hours. You could get decks of cards for statistical routines and for various engineering calculations, attach your data cards, feed the whole pile into a card reader, and return later to the computer center for your printouts, often far into the night when you were trying for as many runs as possible. The programs that I wrote were mostly for numerical solutions of equations. I devised a game that taught my students in biochemical engineering a little about bioprocess development [3]. The punch cards had 72 spaces (fields), so I decided upon 7 variables (sugar concentration, amount of oil, percentage of inoculum, etc.) that each took 10 spaces.

The minicomputer caused a revolution in attitudes. For the first time, the ordinary user could sit at the computer and work interactively with programs. Paper tape replaced punch cards, and magnetic storage devices soon took over. Digital Equipment Corporation sold minicomputers such as their PDP-8 that was inexpensive enough for a few people to share. There was one just down the hall from my office, and I could use it for 4 or 5 h each week. Memory was limited, and programming was at the processor level. You had to code each operation. For example, multiplication required moving binary numbers in and out of the central processor, shifting bits, and keeping track of memory locations. Working with floating point numbers with some bits for the characteristic and others for the mantissa was not easy. You learned to think in

binary and then in octal because it was less cumbersome. Before long there were languages that could simplify operations at an assembly level. Just about the time I learned one, higher level minicomputer languages appeared soon to be followed by compilers for real languages such as FORTRAN. Now you could write code easily, debug interactively, and perform what-if experiments with your programs. Coils of paper tape for storing programs were superseded by flat-fold paper tape. Very tough plastic tape was used to some extent.

Minicomputers made it practicable to dedicate a computer to a process. Groups such as that led by Humphrey at the University of Pennsylvania developed ways to interface a computer to a bioreactor. Numerous students wrote new code or improved the code of other students. Much was learned about sensors, signal conditioning, data display, and process analysis. The concepts were the bases for commercial software, but the code from the early days is mostly obsolete. That is not to say that some groups do not still write code for computer interfacing, but chances are that commercial software will handle most tasks [4]. Instead of a year or more for writing your own program, learning to use commercial software takes perhaps 2–6 weeks.

Personal computers intruded on the monopoly of minicomputers, and you could own a computer instead of sharing with others. The first magnetic storage that was affordable was an audio tape cassette recorder; the stream of bits from the computer produced sounds that could be played back and reconverted to bits. A program might be saved as three or four different files to have high probability that at least one copy would function properly.

My first personal computer, an Altair, was build from a kit in 1976 and had 12 kilobytes of memory. A short program had to be toggled in with switches on the console before the computer could read from a paper tape. You tended to leave your computer on overnight because mistakes were common when toggling, and it could be highly annoying to get it booted again. The version of BASIC that I used took more than 8 kilobytes of the 8-bit memory, leaving little for the code written by me. One inexpensive way to add memory 4 kilobytes at a time was to wire a kit for a circuit board, insert memory chips, and plug the board into the computer.

I must express deep gratitude to students who worked part-time in my laboratory. We usually had a student from electrical engineering who could build devices and troubleshoot problems. Today, all of us can be frustrated when installing new hardware or a new program because the instructions are not always clear and because following the instructions is no guarantee that the results will be satisfactory. This is a picnic compared to debugging problems in the early days. With our home-built computers it was essential to trace circuits, identify bad chips, and to match cables to the ports. When we had better PCs, these electrical engineering students were still of great value for constructing sensor circuits, matching impedances, fixing the A/D converters, connecting stepping motors, and the like. We built our own preamplifiers for \$10 worth of parts, and they performed as well as units costing between \$500 and \$1000. My students complained about taking time to construct and test electronic circuits, but I met students at other universities who complained about equivalent electronic devices that they purchased. There are delays in shipping and lost

time for service with commercial equipment. When something went wrong with a home-made circuit, we fixed it in a matter of hours instead of waiting for days or weeks to get outside service. My students learned enough simple electronics to impress the other graduate students in chemical engineering.

An early input/output device was the teletype. It combined a typewriter, printer, and paper tape punch/reader. Service with a computer was demanding, and repairs were frequent. I recall being responsible for three primitive PCs that were used by students. Each had a teletype, and few weeks went by without lugging one teletype out to my car and going off to get it fixed. Dot matrix printers made the teletype obsolete. These first printers were noisy, and enclosures to deaden their sound were popular. Cost of a printer for your PC approached \$1000, and performance was much inferior to units that cost \$150 today. I have owned dot matrix printers, a dot matrix printer with colored ribbons, a laser printer, and most recently an ink jet color printer that eats up ink cartridges too quickly.

My next personal computer was similar to the Altair, but with read-only memory to get it booted and an eight-inch floppy disk drive. There was some software for crude word processing. Much of the good software came from amateurs and was distributed by computer clubs or could be found at universities. Several years passed before we had graphics capability. I started computing from home by connecting through the phone lines to the university computer center with a dumb terminal. My wife was taking a course in computing, and we had to drive to the computer center to pick up print outs. Our modem was so slow that there was hesitation as each character was typed. A dot matrix printer was soon connected to the spare port on our dumb terminal, and not so many trips to the computer center were needed. Another computer purchased for home used our dumb terminal for display and led to mostly local computing, with the university center available when needed. As faster modems became available, we upgraded for better service. By about 1982, I was using electronic communication to colleagues at other institutions. Software was becoming available for entertainment that provided breaks from serious programming. My wife became a publisher because my books were integrated with teaching programs on a disk, and major publishers were leery about distributing disks and providing customer support for the programs. The university now had a laser printer that we used to make camera-ready copy for my books. My wife learned to use some packages for preparing manuscripts and eventually found that LaTeX was wonderful. The LaTeX commands for spacing terms in an equation are complicated, and I remember how she spent hours getting one messy equation to print correctly.

The Apple computer with full color display when connected to a television set showed what a personal computer could be. Its popularity encouraged competition that brought the price of crude home computers to as low as \$100. Some people in the sciences and in engineering used the Apple computer professionally, but it was not quite right. It was clumsy for editing text because letters large enough to read on a TV screen required truncating the line to only 40 characters. You were better off connecting your computer to a monitor with good, readable, full lines of text. The early IBM computers and the many clones

that were soon available had only a monochrome display, but the monitors were easy to read.

BASIC can do just about anything and is nicely suited to personal computers. It has ways to get signals from a port and to send signals back. Early FORTRAN for personal computers did not come with easy ways for reading and writing to the ports. When most programs were small, it did not matter so much that BASIC was slow. Its interpretative code runs right away, and FORTRAN and the other powerful languages require a compiling step.

Interaction with the computer was at the command line at which you typed your instruction. The graphical user interface was popularized by Apple Computers and was a sensation with the monochrome Macintosh. While the Apple company kept close control of its system, IBM used the DOS operating system that made Bill Gates a billionaire. This was an open system that led to many companies competing to provide software. Apple has done well in some niches for software, but PCs that developed from the IBM system have a richer array of software that has driven them to a predominant share of the market.

I went a different route in the early 1980s with the Commodore Amiga, a truly magnificent machine that was badly marketed. The Amiga was fast and great for color graphics because it had specialized chips to assist the central processor. It had both a command line interface and icons. At one time, I had five Amiga computers at home, in my office, and in the laboratory. I used the command line perhaps a little more often than I clicked on an icon. With today's Windows, it is not worth the trouble of opening a DOS window so that you can use a command line and wildcards to make file transfers easy. The Amiga had true multitasking. This required about 250 kilobytes of memory in contrast to today's multitasking systems that gobble memory and require about 80 megabytes of your hard drive. My first Amiga crashed a lot, but later models did not. My computer purchased in 1998 has the Windows operating system and crashes two or three times each week.

Minicomputers evolved into workstations and developed side-by-side with personal computers. Magnetic storage started with large drums or disks and became smaller in size, larger in capacity, and lower in price. Persistent memory chips stored programs to get the computer up and running. Eight-inch floppy disks were rendered obsolete by 5-1/4-inch floppies that gave way to the 3-1/2-inch disks. The first PCs with hard drives had only 10 megabytes. My first Amiga with a hard drive (70 megabytes) made dismaying noises as it booted. Inexpensive personal computers now have options of multi-gigabyte hard drives. I find essential a Zip drive with 100 megabytes of removable storage. There are devices with much more removable storage, but I find it easier to keep track of files when the disk does not hold too many.

It was a logical step to use the ability of the computer as the basis for word processing. With the early programs, you could only insert and delete on a line-by-line basis. The next advance was imbedded commands that controlled the printed page. I was served very well for about seven years by TeX and its offshoot LaTeX that had a preview program to show what your pages would look like. What-you-see-is-what-you-get seems so unremarkable now, but it revolutionized word processing. The version of LaTeX for the Amiga came with

over a dozen disks with fonts, but there were very few types. These were bit-mapped fonts, and each size and each style required a different file on the disk. I obtained fonts at computer shows, bought some Adobe fonts, and found others in archives at universities. These were intended for PCs, but the files were recognized by my Amiga computer. I had to install them on my hard drive and learned how to send them to the printer. Proportional fonts that are scaled by equations have made my a huge collection of bit-mapped fonts obsolete. There was also incompatibility between PostScript and other printers, but conversion programs solved this problem.

It may seem extraneous to focus so much on the hardware and software, but your use of a tool depends on its capabilities. New users today cut their teeth on word processing, perhaps as part of their e-mail, but this was NOT a common use of computers in the early days. There were few CRT displays except at the computer center itself, and users worked with printed pages of output that were often just long listings of the programs for debugging. These were big pages, and printing on letter-size paper seems not to have occurred to anyone.

Many of us realized that pictures are better than words and wrote programs that showed our students not only columns of numbers but also pages with Xs, Os, and other characters positioned as a graph on the print out. Better graphs were available from a line printer, but there were few of these, and it was troublesome to walk some distance to get your results. There was usually a charge associated with using a line printer, and someone had to make sure that its pens had ink and were working. There is a great difference between computer output as printed lines of alphanumeric characters and output as drawings and graphs. It was quite some time before the affordable small printers for personal computers had graphics capability, but monitors for graphics became common. Furthermore, the modern computer can update and animate its images for its CRT display. BASIC for our computers had powerful graphics calls that were easy to learn. The professional programmers used languages such as C for high-speed graphics. Programs for word processing were followed by spreadsheets and other business programs. With the advent of games, the software industry took off.

3 Biotechnology

Portions of this historical review pertain to academic computing in general, but there were some specific features for biotechnology. Three interrelated areas of particular importance are simulation, process monitoring, and process analysis.

3.1 Simulation

Simulation, an important tool for biotechnology, is considered essential by many bioprocess engineers for designing good control [5]. As you gain understanding of a system, you can express relationships as equations. If the solution of the equations agrees well with information from the real system, you have

some confirmation (but not proof) that your understanding has value. Poor agreement means that there are gaps in your knowledge. Formulating equations and constructing a model force you to view your system in new ways that stimulate new ideas.

Modeling of bioprocesses had explosive growth because the interaction of biology and mathematics excited biochemical engineers. Models addressed mass transfer, growth and biochemistry, physical chemical equilibria, and various combinations of each of these.

It becomes impossible to write simple equations when an accumulation of factors affects time behavior, but we can develop differential equations with terms for important factors. These equations can be solved simultaneously by numerical techniques to model behavior in time. In other words, we can reduce a system to its components and formulate mass balances and rate equations that integrate to overall behavior.

The concept of a limiting nutrient is essential to understanding biological processes. The nutrient in short supply relative to the others will be exhausted first and will thus limit cellular growth. The other ingredients may play various roles such as exhibiting toxicity or promoting cellular activities, but there will not be an acute shortage to restrict growth as in the case of the limiting nutrient becoming exhausted.

The Monod equation deserves special comment. It is but one proposal for relating specific growth rate coefficient to concentration of growth-limiting nutrient, but the other proposals seldom see the light of day. This equation is:

$$\mu = \frac{\hat{\mu}S}{K_s + S} \quad (1)$$

where

μ = specific growth rate coefficient, time⁻¹,

$\hat{\mu}$ = maximum specific growth rate, time⁻¹,

S = concentration of limiting nutrient, mass/volume, and

K_s = half-saturation coefficient, mass/volume.

Students in biochemical engineering tend to revere the Monod equation, but practicing engineers apply it with difficulty. There is no time-dependency; it is not a dynamic relationship and cannot handle sudden changes. Industrial batch processes encounter variations in the characteristics of the organisms during the run such that coefficients on the Monod equation must be readjusted.

Simulation paid off. One of my students, Thomas Young, joined Squibb in about 1970 and soon made major improvements in the yields of two different antibiotic production batches, mostly as the result of simulation. I had recommended Tom to my old employer. They declined to make him an offer because they considered him too much of a theoretical type. A vice-president at Squibb told me that Tom was just about the best person that they ever hired and that his development research saved their company many millions of dollars. It was partly the ability to test ideas on the computer that led to rapid progress, but even more important was the thought process. Deriving equations for simulation forces you to think deeply and analytically, and many new insights arise.

3.2

Monitoring and Control of Bioprocesses

Instrumentation in a chemical plant brings to mind the control room of a petroleum refinery with its walls lined with a cartoon representation of the processes with dials, charts, controllers, and displays imbedded at the appropriate locations. Operators in the control room observe the data and can adjust flow rates and conditions. Such control rooms are very expensive but are still popular.

There are hybrids of the traditional control room with computers for monitoring and control. One computer monitor has too little area for displaying all of a complicated process, and the big boards along the walls may still be worthwhile. However, smaller processes can be handled nicely on one screen. Furthermore, a display can go from one process unit to another. A good example is the monitoring of a deck that has many bioreactors. One display can be directed to a selected reactor to show its status, often with graphs showing variables vs time. The operator can choose the reactor of interest, or each one can come up for display in sequence. Logic can assign high priority to a reactor that demands attention so that it is displayed while an alarm sounds or a light flashes. If we agree that an operator can only watch a relatively small amount of information anyway, it makes sense to conserve financial resources by omitting the panel boards and using several computer displays instead. There is the further advantage that the computer information can be viewed at several locations; some can be miles away from the factory. Connecting from home can save a trip to the plant by a technical person at night or at the weekend.

I think that bioprocess companies have done well in using computers to monitor their processes. The control rooms are full of computers, but the adjustments tend to be out in the plant. Only recently have plant operations been fully automated or converted to control by sending signals from the computers. Because hardware and the labor to set it up and wire the connections have a daunting cost, the lifetime of a computerized system is roughly 5 years. Advances in the technology are so rapid that a system after 5 years is so obsolete that it is easy to convince management to replace it.

I have an anecdotal report of early attempts at automation. The Commercial Solvents Corporation in the 1950s attempted to automate sterilization of a reactor. There was no computer, and the method relied on timers and relays. Unreliability due to sticky valves and faulty switching resulted in failure. Being too far ahead of their times gave automation a bad name. Development engineers at other companies who proposed automation were told that it had been tried and did not work. Today, there is nothing remarkable about computerized bioreactors and protocols for their operation. I have observed automated systems in industry that are fully satisfactory. The computer makes up the medium and can control either batch or continuous sterilization.

3.3

Bioprocess Analysis and Design

There is not a great difference now from what was being done in the past, but there have been many changes in convenience and in capabilities. Engineering professors wrote programs that assisted process design with such features as approximating physical properties from thermodynamic equations. These properties are crucial to such tasks as designing distillation columns but do not matter much in biochemical engineering. Today, there are excellent commercial programs such as Aspen that will develop the required thermodynamic properties en route to designing a process step or even an entire system. I experimented with Aspen and my opinion of it comes later.

4

Recent Activities

A major advance has been databases and programs that manage databases. Libraries of genetic sequences have become essential to many biotechnologists, but this area deserves its own review and will not be mentioned again here.

4.1

Models

Models should be judged on how well they meet some objective. A model that fails to match a real system can be highly valuable by provoking original ideas and new departures. Overly complicated computer models, of course, can have a fatal weakness in the estimation of coefficients. Coefficients measured in simple systems are seldom valid for complex systems. Often, most of the coefficients in a model represent educated guesses and may be way off. Complicated models take years to develop and may be impractical to verify. Such models are worth something because of the organized approach to just about all aspects of some real system, but there are so many uncertainties and so many opportunities to overlook significant interactions that predictions based on the models may be entirely wrong.

Deterministic models (those based on actual mechanisms) make a great deal of sense when they are not too unwieldy. The terms have physical or biological meaning and thinking about them may lead to excellent research. The goal of the modeler should be to identify the most important effects and to eliminate as many as possible of the insignificant terms. It always comes back to the purpose of modeling. To organize information, we may just keep adding terms to a model in order to have everything in one place. When the goal is prediction, a model should be tractable and reliable. That usually means that it must be simple enough that its coefficients can be estimated and that the model can be verified by comparing its predictions with known data.

Most real-world situations are too complex for straightforward deterministic models. Fortunately, there are methods that empirically fit data with mathematical functions that represent our systems and permit comparisons and predictions.

Models that assume steady state are usually fully satisfactory for continuous processes. Continuous cultivation tends to steady state or may fluctuate fairly slowly so that the Monod equation can be applied. Batch processes and dynamic situations in continuous processes must consider non-steady states. Franzen et al. [6] developed a control system for respiratory quotient and found that steady-state models were unsatisfactory.

The term *hybrid* brings to mind hybrid computers that combined an analog computer and a digital computer. The term *hybrid model* is applied when a model has components that attack a problem quite differently. For example, there might be deterministic features, aspects of neural networks, and optimization routines that must be organized and integrated [7, 8].

Computer models aid state estimation for bioprocesses. Such models have great value for predicting the best harvest time for a bioreactor when the product concentration peaks and declines or when the rate of product formation no longer repays the cost of continuing the run.

4.1.1

Unstructured Models

The simplest models lump relationships. A good example is letting X stand for cell mass. This is an absurd oversimplification because cell mass is composed of hundreds of biochemicals that can change in proportions and that function very differently. Nevertheless, growth dynamics may be represented fairly well by the simple equation:

$$\frac{dX}{dt} = \mu X \quad (2)$$

Letting S stand for the concentration of growth-limiting nutrient is another example of lumping because a real nutrient medium can have several ingredients that contribute. For example, the intermediary metabolism of fats produces compounds for carbon-energy, but we may focus on a sugar as the limiting nutrient.

4.1.2

Structured Models

Ramkrishna's group at Purdue University uses the term *cybernetic* for biological models that interpret the control actions of organisms [9, 10]. Current interest seems to be focused on simultaneous or preferential uptake of carbon sources. By incorporating cybernetic concepts, they can predict diauxic growth patterns as well as simultaneous use. The structured model considers precursor compounds and the patterns for their formation and consumption.

Structured models can become complex when the fine details of growth, biochemistry, and equilibria are incorporated. One example is a description of *Escherichia coli* developed under the leadership of Shuler at Cornell University [11] that has dozens of biological effects, a multitude of chemical reactions, and hundreds of coefficients.

A fungal bioprocess modeled in great detail qualifies as a structured model even though some aspects of the biology are lumped [12]. For example, fungal morphology (such as pellet size and the fraction of pellets in the total biomass) is a structural feature that could be further categorized for the biochemical details.

4.2

Bioprocess Control and Automation

Appendix 1 has some terminology of process dynamics and control for readers who are not engineers. The theory of process control stays ahead of practice, and some aspects of the theory have minor importance for biotechnology because our processes change slowly and have no analogy to exothermic chemical processes that can explode. There are still times when it makes sense to purchase a conventional controller with proportional-integral-derivative control. For example, a storage tank may benefit from temperature control, and an inexpensive controller that adjusts the flow of cooling or heating water through a jacket or coil may suffice. However, most factories have computers that control their bioprocesses, and there are enough places to hook up sensors and to send signals to controllers to accommodate some more control. When you run out of channels, you buy another interface board or another computer. The cost can be far less than purchasing several controllers, and the entire control system can be displayed conveniently in one location instead of at the individual controllers that may be widely scattered. Sonnleitner [13] provided perspective on automation and design of bioprocesses and addressed some potential pitfalls. Galvanauskas et al. [14, 15] have a simple procedure for enhancing bioprocess performance through model-based design.

The concepts of proportional, integral, and derivative control retain their importance, but using a computer in place of a controller permits far more sophistication [16]. The settings for the gain of the controller and its coefficients for its various actions are compromises because the process can change. This is particularly true of bioprocesses. Batch processes start with high nutrient concentrations, few organisms, and distinct properties such as temperature, pH, viscosity, and the like. As the run progresses, concentrations change as do physical properties. The controller settings that worked well early in a run may be poor at a later time. The control algorithm in the computer will employ concepts of proportional, integral, and derivative control, but can augment them with logic. In the simplest case, past experience may be used to devise a schedule for changing the control coefficients in steps, e.g., when the time reaches some point, and switch to these coefficients. In the more advanced cases, a model of the process makes the decisions for instantaneous adjustments of the control coefficients.

4.2.1

Sensors

Manual sampling at frequent intervals was the norm for industrial bioprocesses for many years. Gradually, more and more methods have been developed for on-line measurements. Until the early 1960s there were no reliable pH electrodes that could be sterilized. Several of our bioprocesses performed best at the correct pH, and we put a great deal of effort into taking samples, titrating, and adjusting the pH in the vessel. The first sterilizable electrodes had short lives. If the pilot plant had 20 vessels with internal pH electrodes, half might fail during a run. Although expensive, an internal pH electrode is standard operating procedure today.

The extensive research and development of better sensors for bioprocesses is beyond the scope of this review [17]. One example of the level of sophistication of current procedures is Rothen et al. [18].

4.2.2

Observers

An observer is a method for mixing models and real data. The general concept is that some variables can be measured by cost-effective analytical procedures while other variables are costly, troublesome, or impractical to measure. The observer uses both the model and the practical measurements to estimate those variables that were not measured directly. An example of something that is impractical to measure continuously is cell mass. The usual estimate of cell mass draws samples, collects the solids by filtration or centrifugation, and dries and weighs them. This is time consuming and laborious, so the value for cell mass is unavailable when needed as a control variable. There are alternative methods of estimating cell mass by optical density, nucleic acid content, or the like, but such methods track imperfectly with cell mass. A good model of the bioprocess will have cell mass as one of its terms. We can construct the model so that it uses the measured variables and estimates cell mass. If the model is reasonably good, we can use the estimate of cell mass as an index to control the process.

Bioprocess variables that are usually measured continuously are pH, temperature, and feed rates. Other possibilities are impeller speed, aeration rate, concentrations of carbon dioxide and oxygen in the exiting gas, and perhaps some concentrations in the medium. Some textbooks refer to the assays that are used for estimation of other parameters as *gateway* sensors because they open the door for using the model.

Models are not perfect representations of a process. As the bioprocess and its model drift apart, estimates made by the model will be incorrect. Errors propagate with time and become more serious. However, we can measure critical variables occasionally and correct the estimates. An analogy would be an internal pH electrode that drifts haphazardly. Before the next run, we would replace the failing electrode with a good electrode, but the bad electrode may be all that we have right now. We can still use its readings if we take samples from the bioprocess, measure the pH accurately at the lab bench, and adjust the reading

of the poor electrode. If the drift is very large, we must recalibrate more often. In any event, the pH signal gives us something for continuous control of the process, and control based on grab samples is a very unattractive alternative. When resetting the pH reading, we should correct to the time that the sample was taken, not to the present time because the electrode may have drifted during the sampling.

The above method is also applied to observers, but there are other ways to correct. One useful technique is to estimate something that can be measured continuously. The estimates are compared with the true values to decide how much the model is in error, and the amount of error determines the compensation for other estimates. This is really just a slight difference from using the additional measurement as part of the model.

Off-line measurements are of limited value for control but can be major assets for checking a model and correcting estimates by an observer. Poor models are risky. If the model is unstable, model errors can give rise to estimation errors that increase exponentially. This is an example of when bad control is worse than no control.

4.2.3

Auxostats

The concept of an auxostat is based on process control. The nutrient feed stream is used to fix some variable of the process. Pumping dilutes concentrations in the medium. If the microbial population is unable to maintain the desired concentration, the pump is automatically slowed down while the organisms both grow and excrete product. Once steady state is established, the specific growth rate of the organisms equals the system dilution rate [19]. I think that Edwards et al. [20] were the first to publish a description of an auxostat, but Watson [21] previously did something quite similar. The turbidostat, first used in the 1940s, has this same concept, but the property being controlled is turbidity of the culture.

Auxostat operation of a continuous reactor system holds the promise that the culture can be operated safely at a high specific growth rate [22]. When, for example, the product is cell mass, dilution by fresh medium is offset by growth. Furthermore, there is intense competition among the organisms. This operating mode has the advantage of washing out the slower growing microorganisms from the reactor leaving more substrate for those that grow faster. It is also obvious that a faster growing population (from mutation or contamination) challenges the established culture to cause its loss from the system. Auto selection of yet faster growing cultures is the logical result. When the specific growth rate should be fixed and not allowed to seek its own maximum, the control method of Levisauskas et al. [23] can be used.

4.2.4

Examples

Some typical examples of computerized control are shown in Table 1.

Table 1. Examples of computer control

Product or Process	Strategy	Reference
Glutathione	Advanced control	24
Acetate	Internal model	25
L-Carnitine	Analysis of model	26
6-Hydroxynicotinic acid	Analysis of model	26
5-Methyl-2-pyrazincarbonic acid	Analysis of model	26
Nicotinamide	Analysis of model	26
Alkaline protease	Better monitoring	27
Acetone/butanol	Metabolic analysis	28
Penicillin	Hybrid model	29
Penicillin enzymatic deacylation	Kinetics, equilibria	30
Lipase	Growth rate control	31
Brewers yeast flocculation	Analysis of model	32
Waste water treatment	Neural network	33

4.2.5

Modeling and Control of Downstream Processing

Some aspects of recovery of products after bioprocessing are relatively straightforward. Admittedly, there are complications due to chemical equilibria and reaction kinetics, but these are not unique to processes for purification of product streams. Downstream processing incorporates the main lines of chemical engineering and can draw on a wealth of experience. Software tools and models were discussed by Simutis et al. [34]. The commercial programs, such as Aspen, that are used to design chemical processes can be used as well for downstream processing, but I have not been impressed very much by their applications to biochemical engineering. One beauty of such programs is handling of physical properties. Essential coefficients can be drawn from a library that is part of the program, or they may be estimated with thermodynamic equations. While this is a great aid for the usual chemical process industries for such steps as distillation, there are not many biochemical engineering steps at high temperatures or pressures where you need to know the thermal properties of the process streams. Practically all of biotechnology is aqueous.

Having stated that there is little difference from regular chemical engineering in terms of the intellectual challenges of computer approaches to downstream processing, I must admit that in almost all cases, the recovery operations cost much more than the bioprocessing. It makes a great deal of sense to improve costs through better modeling and control of downstream processing. Bulmer et al. [35] have applied computer simulation for improving the recovery of intracellular enzymes and demonstrated this with alcohol dehydrogenase from *Saccharomyces cerevisiae*. Zhou et al. [36] modeled the interrelationships of bioprocessing and product recovery to optimize both. Vanhalsema et al. [37] modeled phase equilibria to improve extraction of carboxylic acids.

4.3 Intelligent Systems

State-of-the-art control systems employ artificial intelligence. Several companies now offer complete systems for monitoring and control of bioprocess, and the control algorithms use advanced logic. Two web sites for further information are <http://www.biospectra.ch> and <http://www.gensym.com>. One of my former students worked for GenSym and played a major role in developing such a system. The following descriptions of current practice are based on pages downloaded from their web site.

Example 1: Gensym's G2 Diagnostic Assistant (GDA) was installed in the fermentation plant at Eli Lilly's Tippecanoe Laboratories (Lafayette, IN) with three goals: control its antibiotic production processes by reducing variability, increase yield while reducing unit costs, and give operators the responsibility for the process. Using statistical information from previous runs, control charts were developed. This system monitors incoming production data, compares them against the established control charts, and provides operators with expert advice. A distributed control system collects and processes data, provides expert instructions in real time, and also provides an intelligent operator interface that advises the operator on how to respond to unusual behavior or informs the operator that the situation is sufficiently novel that no advice can be offered.

In the first month of system operation, process deviations dropped 72.3% and yields increased 3.2%. As a result, the company has realized significant cost savings and has become better able to match production to changes in sales.

Example 2: The Bioindustrial Group at Novo Nordisk (Denmark) uses a Gensym system for monitoring, sensor validation, diagnostics, and state estimation of fermentation processes for the laboratory scale production of industrial enzymes produced by genetically engineered microorganisms. Suitable sensors may not be available to determine the metabolic state of the processes, so the system combines sensor data with analytical data for state estimation of the culture's metabolic status. Bioprocesses have many problems with sensors. This system continuously validates sensors using statistical calculations and mass balances and identifies deviations from normal or expected behavior while providing high level supervisory control. Controller "responsiveness" is an important aspect of Novo applications, and the system makes the control more responsive to variations in the culture using empirical on-line optimization.

Stephanopoulos and Han [38] have reviewed intelligent systems in process engineering. Quite involved or complicated logic can be programmed. Instead of fixed setpoints or controller settings, there can be a schedule for adjustments, or logic can decide the settings. Furthermore, artificial intelligence can analyze the process to establish the control specifications. Diaz et al. [39, 40] have used predictive control of the concentration of dissolved oxygen in a bioreactor.

4.3.1

Expert Systems

Control can be based on rules in an expert system. For example, an expert system can take into account the lag phase and the youthful physiology of the cells for control of the early stage of a bioprocess while the control can be much different for other phases. Programming simple rules as IF/THEN statements is not a serious challenge for control engineers, but commercial programs called *Expert System Shells* can usually do it better. These shells provide for compound rules, rule priorities, exceptions, explanations of why certain rules have fired, and more. One important report is the rule activity chart that tells which rules fired, when they fired, and how intense was their action. This helps greatly when revising and improving rules. Furthermore, comparing rule activity of one run with another run can improve understanding of the process and the ways for improvement.

De Bernardez et al. [41] developed an early expert system for controlling a bioprocess. It was particularly interesting that the expert rules had to be selected carefully to avoid exhausting the available memory of the computer. This seems archaic now when personal computers have enormous sizes of memory, but not long ago memory could seriously limit what could be done. The maxim “work expands to fill the available time” has the analogy that computation can expand to tax the available memory. For example, collecting data from several bioprocesses at frequent intervals can create gigantic files. Say that the sampling rate is 100 times per second and that a run lasts for 100 h. If there were 10 sensors to be sampled and 10 calculated points to be plotted, the file size to hold all this information would be:

$$100 \text{ h} \times 60 \text{ min/h} \times 60 \text{ s/m} \times 100 \text{ times/s} \times 10 \text{ sensors} \times 10 \text{ calculations} = 3.6 \times 10^9$$

It is not wise to use memory resources this way; the data should be averaged for a time interval. For example, the average pH during 1 min of the run may be adequate with no need for knowing its value each one hundredth of a second. If the file is still too large, it should be compressed.

4.3.2

Fuzzy Logic

One organized method for dealing with imprecise data is called *fuzzy logic*. Even with precise information, real-world problems may include uncertainty. There is a range of values for failure, and the designer of a bioreactor will incorporate safety factors to make collapse or rupture highly unlikely. Unreliable data magnify the chances for erroneous conclusions, and fuzzy algorithms commonly estimate the level of confidence of the results. The data are considered as *fuzzy sets*. Traditional sets include or do not include an individual element; there is no other case than true or false. Fuzzy sets allow partial membership. These memberships can be manipulated with the operators shown in Table 2.

As would be expected, probability is often needed to clarify fuzzy, uncertain information. A degree of truth is the probability that a statement is true. For a

Table 2. Fuzzy operations

Operation	Definition
Fuzzy AND (f-AND)	a f-AND b = $\min(a, b)$
Fuzzy OR (f-OR)	a f-OR b = $\max(a, b)$
Probability AND (p-AND)	a p-AND b = $a * b$
Probability OR (p-OR)	a p-OR b = $a b - (a * b)$
(same as Unary NOT)	NOT a = $1 - a$

practical situation, we might use one of the following methods to design a fuzzy controller [42]:

1. Base the controller on a human operator's experience and/or a control engineer's knowledge
2. Model the control actions of a human operator
3. Base control on a fuzzy model of the process

The first method is analogous to constructing an expert system, and this is the most common approach to fuzzy control. The first and second methods can only approach the performance of a human because they strive to mimic a human. Some processes are so complicated that no humans can control them well. For example, the U.S. Air Force has a fighter plane that cannot be flown by hand – the computer must fly it with the human supplying information about where to go and how fast. To go beyond human capability, Method 3 with a fuzzy model of the process has the best chances for success. The problems are to identify the system features in a form that may be modeled and to design an effective controller that can be applied to this model.

Fuzzy reasoning can incorporate heuristic knowledge into the model. Simutis et al. [43] report using fuzzy information with an extended Kalman filter to follow alcohol formation in beer brewing. Their model outperformed conventional approaches for estimating the process state of an industrial scale reactor. Their linguistic variables had the following forms:

- Process time: short, medium, long
- Sugar concentration: low, medium, high
- Process temperature: low, medium, high
- Sum of evolved CO₂: small, medium, high
- CO₂ evolution rate: low, medium, high

Truth tables were developed to assign membership in these categories. Precision of some of the assay methods was low, and the fuzzy decisions were more appropriate than using the numerical values. More recent advances incorporate state modeling with fuzzy control [44].

4.3.3

Neural Networks

Neural networks represent knowledge in an adaptive, distributed architecture [45]. The general idea is relating outputs and inputs in a manner akin to *pattern*

recognition. There are input nodes and output nodes with one or more layers of nodes in between. Weighting coefficients for the summation of inputs to each node develop the outputs. The inputs and outputs for sample cases are used to improve coefficients through iteration, and eventually the system can derive reasonably correct outputs from inputs never before encountered.

A crude example of how a neural network could be useful is a process with several different feed streams and measurements such as concentration, temperature, and pH. There will be permutations of our input information for which the output is known. Output may be action rules such as add more acid, increase the rate of cooling water, sound an alarm, or the like. As our neural network iterates and learns with solid examples of appropriate outputs for given sets of inputs, it can evolve until it recognizes patterns in these inputs. When presented with totally new inputs, it may be able to generate a correct output. The correctness of the output will depend on how well it has been trained, but insufficient examples from which to learn will give unreliable answers.

Important facts about neural network programs are that learning is slow (medium-sized networks may take hours of training on a slow computer) but decisions with a trained network can be lightning fast. The learning requires iteration, error checking, and testing for convergence. Calculating the output of a trained network is merely once through with multiplication and addition. Commercial programs called *shells* are easy to use with no mathematics. Furthermore, the neural network learns automatically; you configure it but do not have to write any program. The neural network tends to be as good as its training data. A well trained and carefully tested network is a highly reliable system that did not need many long hours of programming by a human.

Vancan et al. [46, 47] have presented strategy for using neural networks in bioengineering situations. Simutis and Lübbert [48] discussed developing process analysis through mass balances and neural networks and augmented such analysis with random search optimization routines that find optimal regimes of feed rate, feed concentration, temperature, or precursor profiles [49].

Stoner et al. [50] have used computer interfacing and control with integrated software packages including learning to deal with multiparametric effects on the oxidation of iron by thermophilic bacteria. Shimizu [51] has combined fuzzy logic with a neural network and provided three examples of processing with *Saccharomyces cerevisiae* that used computerized state recognition and fuzzy control of the processes.

4.4

Responses of Microbial Process

Steady state analysis is becoming outdated because there is no steady state in a batch process while the organisms are alive, and control of continuous processes must consider dynamics to be effective. A report that it took time for microorganisms to respond to forcing by Mateles et al. [52] affected my thinking very nearly at the start of my professional career, but a decade later there were still grievous errors in dynamic analysis. The same Thomas Young that I lauded earlier presented a paper at a national meeting of the American Chemical

Society in about 1970 and upset the audience by criticizing the Monod equation. It seems so obvious now that this important equation has no time dependency. The specific growth rate cannot respond instantaneously, so the equation is not equipped to handle abrupt changes in nutrient concentration. Nevertheless, it was heresy at that time to criticize one of the tenets of biochemical engineering.

With no delay in growth-rate adjustment, the response of a bioprocess to an upset should move smoothly to the new final value without overshoot. Real bioprocesses tend to overshoot because organisms require time to adjust their biochemistry to new environmental factors.

Most university computer centers have programs that aid in dynamic analysis, and some programs for PCs have appeared. The user can specify the scheme of a system (this is termed the *block diagram*) and the time constants. Either the connections of blocks or the coefficients can be changed to determine the response and stability. There are also features for analyzing stability.

4.5

Metabolic Engineering

Metabolism and biosynthesis depend on parallel and sequential biochemical reactions subject to controls based on induction, activation, or inhibition of the required enzymes. Intermediate biochemicals progress through competing pathways at rates determined by the activities of these enzymes. The manipulation of these enzyme activities is termed *metabolic engineering*. Some control is possible by adding precursor compounds or substances that affect enzyme synthesis. For example, a number of carbohydrates induce or stimulate the formation of cellulase enzymes. Adding them will result in higher activity of cellulases. However, most metabolic engineering comes from genetic alterations to cripple certain unwanted pathways or to promote desired pathways. Computer models of the reactions and biochemical controls aid research by highlighting the enzyme activities of interest and predicting the effects of changes.

Metabolic engineering is one of the most exciting areas of biotechnological research but is beyond the scope of this review. A few typical references can be consulted [53–57].

5

Information Management

My years in industrial process development were in the dark ages of data processing. We collected hand-written records kept by the operators of the results of samples taken for pH, turbidity, packed cell volume in a centrifuge tube, temperature, amount of antifoam added, concentration of product, and the like. We requested assays for sugar concentration, amino acids, and specific biochemicals when appropriate. Everything was on paper and circular charts or strip charts. To compare one run with another, you had your assistant and maybe other people covering one or two tables with paper, sifting through, and

calling out information from one run as you inspected another. This was such a chore that we did a rather poor job of tracking the history of runs to learn how the best runs differed from the others.

With a computer interfaced to a bioprocess, the collected data are already in a format suited for a spreadsheet or database program. These programs have features for displaying results as graphs or charts. Commercial database programs make it easy to compare different experiments and runs. These programs include routines that manipulate and treat the data. We can easily set graphs side-by-side or even superimpose them. There is no excuse for not searching for reasons why some runs are better than others. Developing better ways to search these databases is an active area of research [58].

Networking of computers ranges from a few stations to the whole world. Laboratory information management systems (LIMS) substitute computer actions for some routine, boring human labor and eliminate errors from entering and transcribing data.

5.1

Customer Service

Some companies have gone far beyond routine advice to customers by phone, fax, e-mail, or personal visits by integrating their computers with those of their customers. A good example is Iogen Limited, based in Ottawa, Canada. They get on-line reports from computers that control customers' reactors that use Iogen enzymes. When there are process deviations, recommendations are made almost immediately. There was an anecdotal report of another company attempting to gain market share from Iogen by offering similar enzymes at one-half the price. The customer declined that offer with contempt, and the manager who had hoped that price alone was the key to sales was fired.

5.2

Electronic Communication and Teaching with Computers

All areas of science and engineering make use of e-mail, fax machines, and the Internet. Information today flows at much faster rates than in the past and in huge amounts. Graphic information benefits from computers. Digital cameras and scanners are inexpensive. We can include color images in our communications very easily. Color printing that was costly just a few years ago is now routine with printers sold in computer stores for as little as \$150.

Biochemical engineering is well-served by web sites that make our lives easier. In preparing this review, it took but a few hours to search for the authors whose research I had been following and for keywords to compile examples of current work. Our computer programs allow cutting and pasting from one to another. I collected references as files from the Internet and cut out the needed information with little retyping.

I strongly support electronic publication, but the emphasis so far is on better distribution. Short times between submission and publication are important advantages of handling manuscripts with e-mail, and placing reports that look

much like the old hard copies on the Internet insures wide and timely distribution. However, electronic journals in their infancy have hardly begun to use modern technology. I think that an exciting development is pages or displays that do something, and this was my main reason for learning Java.

Science and engineering are sprinkled with equations. Graphs based on equations are appreciated and convey information more effectively than just presenting the equations and their terms and symbols. Books and articles cannot afford very many figures, but computer code has no limitations on the number of graphs it draws. By programming to draw graphs of an equation based on reader input, students can explore permutations of its coefficients, learn quickly, and retain a feel for which terms matter most and how sensitive is the result to each coefficient. My educational projects have led to web pages that can be accessed from all over the world. Having my Java code as a template, I can create an interactive web page with a different equation in 20 min. I have extended this to simulations of microbial processes for which the reader can manipulate the parameters.

Examples are applets with scrollbars that change the coefficients for graphing of normal distributions, the Monod equation, adsorption with either the Langmuir equation or the Freundlich equation, Michaelis-Menten enzymatic kinetics, and various growth rate expressions for multiple nutrient limitations. Several of my applets mimic an experiment. The reader selects some conditions, and the computer displays the expected results. Other portions of the page can suggest things to try and what to observe. There can be links to explanatory material. Some of my applets show solutions of simultaneous ordinary differential equations such as dynamic responses to various forcing functions and interactions of microorganisms.

Games grab student interest. By keeping alert to opportunities for teaching with a game, I have had about 5 successes in 38 years. You have to decide what you are trying to teach before you devise some complicated game. An approach that has worked for me is to develop some sort of role playing and to award hypothetical profits that depend on how well you understand your role. Students appreciate money and enjoy making even simulated profits.

Electronic documents can have all sorts of motion, even video if you are willing to wait while it downloads. Sound with the video is seldom important for biotechnology, but a voice could add a personal touch. You do not need Java to present animations. A number of methods depend on shuffling through a collection of images to make a movie. Java also has features for doing this. When simple shapes will suffice for figures, Java subroutines can easily move them around the screen for what I call *algorithmic animation*. One of my animations that required 1 megabyte for separate images was replaced by about 5 kilobytes of Java code that sketched the same moving images (a 200-fold saving). Since this code runs fast right at your local computer, you wait only a moment to download the small Java applet before the animation starts. My Java applets can be found at <http://www.rpi.edu/dept/chem-eng/biotech-enviro/biojava.htm>

6 Some Personal Tips

Modern bioprocesses are seldom operated in strictly batch mode where nothing but air is added after inoculation. Certainly for research there will be some additions of nutrients, reagents, or special biochemicals. Such additions require pumps, and small, reliable pumps are not inexpensive. An adjustable pumping rate is more desirable than a fixed rate, but variable rates pose problems for the computer. Some very nice pumps with variable speed motors are commercially available. These require special control boards that plug into the computer or the interface may connect to one of the ports of the computer. The computer sends one signal to the pump to set its speed and another signal to turn the pump on or off. We have had excellent experiences with these adjustable pumps except that their cost is relatively high.

Constant speed pumps are less expensive than variable pumps, but the rate may be matched poorly to the need. A brief interval of pumping may overshoot the set point and upset the process. We have solved this by establishing a pumping cycle of the order of 10–20 s, an interval that appears nearly continuous on the time scale of microbial processes. Our computer turns a constant speed pump on for a fraction of this interval. For example, to pump at half the maximum rate of the pump, it would be turned on for 50% of the cycle. The electronics for doing this cost very little, and the parallel port of a computer may have sufficient power for the switching. If not, one transistor can develop the needed power. An optical relay on the computer side of the circuit isolates the pumping circuit. When the computer turns on a tiny light-emitting diode in the relay, voltage is applied to the pump. These relays cost from \$1 to \$10 depending on how much current must flow. Heating circuits can use this same principle and at far lower cost than analog controllers of high wattage that vary the voltage to a heater.

Stepping motors are another way to get variable speed. Each pulse sent to the motor control circuit turns the motor a few degrees. The motor speed can be geared down to get a small fraction of a degree per pulse. This is a superb method for getting variable speed for low torque situations, but heavy-duty stepping motors are expensive. Stepping motors for light duty are ubiquitous for moving paper through printers and are a quite inexpensive. We have enjoyed the excellent performance of stepping motors connected to personal computers to drive home-built micromanipulators for our research with microelectrodes inserted specific distances into microbial films. We used an O-ring as the belt to link the stepping motor to the fine adjustment of a microscope body to which our microprobe was fastened. Each pulse from the computer raised or lowered the microprobe by a fraction of a micrometer. The programming to send pulses to the parallel port of a personal computer is simple, and one integrated circuit chip (cost about \$8) adapts pulses to the wiring for the stepping motor.

One daunting feature of interfacing a computer to a bioreactor is obsolescence. A good workstation is expensive, but computers evolve so rapidly that a new model that can run the improved software is needed within two or three years. A computer purchased for several thousand dollars a few years ago

is worth only a few hundred dollars today. However, we have found that old computers and their old data acquisition systems can be devoted to an individual bioreactor. Obsolete computers that were destined for the scrap heap serve in a master/slave arrangement. The master computer should be modern, but the slaves merely have to collect data and pass instructions from the master computer on to the control system of the bioreactor. When a slave computer needs repairing, we return it to the junk pile and replace it with another old computer. The analog-to-digital boards in old computers also wear out, but usually they fail one channel at a time. We move the wires from the sensors to a spare channel and continue on.

7

Conclusions and Predictions

Wieser and Satyro [59] have described current computing practices of chemical engineers and predicted the future. The bioprocess industries and their support network with universities and governmental organizations have benefited greatly from computers. You expect to see a computer at the desk of every scientist and engineer, and these computers are commonly networked. A period of each day is devoted to e-mail. The world wide web is used for retrieval of information. Our journals are moving to the Internet. Many bioprocesses are controlled by computers. When you have the proper permissions and are on the same network, you can look over the shoulder of the operators and follow a bioprocess in real time with your personal computer. It is easy to compare data from different runs, and there are software packages to aid your analysis.

I think that one feature that is lagging is “pages that do something.” We are barely scratching the surface for better communication by adding interactive features to our documents. I am trying to do this with educational presentations, but the concept can be extended to all communications. Instead of long narrative documents, we can hyperlink from an index or abstract. The reader need not plow through the narration but can select the material of most interest. There can be links to supporting material, to spreadsheets, to archives of explanatory topics, to sketches, animations, video clips, and the like.

Programming skills of the typical student have deteriorated. Much of this can be blamed on commercial programs that render many tasks so easy. Our students can be expert with a word processing program, with spreadsheets, and with mathematical aids such as Maple or Matlab. They tend to master new programs quickly. However, they seem reluctant to write computer code. They would rather program their pocket calculator than write code. I have seen students write routines of great complexity using a spreadsheet program, never realizing that it would be much easier to write a program in BASIC or FORTRAN. I excuse this because modern spreadsheets have menu-driven schemes for developing spectacular graphs and charts. I have written code for so many years that I can get a useful graph almost as quickly in my own programs. The situation will probably get worse as the path of least resistance is to muddle through with the commercial programs that you know best.

Much of the exciting research of today will continue for several decades with incremental improvements and refinements. There are some areas, however, when current knowledge is meager and young investigators can make their marks. One is biological control mechanisms. There are some proposals for equations to handle mixed substrate systems, mixed culture systems, toxic nutrients or ingredients, preferential use of nutrients (diauxie), and the like. However, very little is known about how cells sense concentrations, how the information is processed, and how the controls function. A good example is the growth rate of cells. The Monod equation has been invaluable for describing what happens, but it says nothing about how the cellular controls develop the specific growth rate.

I predict that computerized presentations will more and more substitute for lectures. In fact, the lecture would be much more effective after preparing the students with a computer presentation. There is an analogy with a medical examination. The physician relies on tests before seeing the patient. A lecturer could perform better with students who have reviewed background material; the lecture could be modified if the teacher knew more about their strengths and weaknesses.

Fortunes have been made by locking users into an operating system. That may soon end as the Internet switches to its own operating system. At present, a file from the Internet must meet the specifications of an operating system. A much more simple operating system could handle display of files from the internet, and languages such as Java could execute faster by controlling the computer themselves instead of having another layer of executions demanded by an operating system. This could also expedite a revolution in the distribution of software. Instead of upgrading software yourself, you will pay for an Internet service that sends you the programs you need when you need them. These services would compete by having the newest and best software. I predict that commercial software will be layered so that you will not have to download all of a gigantic program. If you invoke a special feature, there might be a brief delay as this program element is downloaded and connected to its master program.

Biotechnologists are always among the first to embrace the advances in computers and software. As this continues, biotechnological education and research will make the bioprocess industries even more of a mainstay for improved living standards and for the global economy.

Appendix: Terminology for Process Dynamics and Control

– Block diagram

The process is visualized as a series of interconnected functional blocks that correspond to process steps or to the controllers or the process sensors. A block acts on its input information and generates output. There may be several incoming signals and several output signals for a block. Typical signals in a bioprocess are concentrations, reaction rates, oxidation-reduction potential, pH, temperature, enzyme activities, and the like.

- Derivative or rate control
Corrective action is based on the rate of change of the error. This is particularly suited to systems that can blow up. If the error is increasing rapidly, a derivative controller can institute a large amount of correction. It is uncommon to use solely derivative control. An industrial controller may combine proportional, integral, and derivative control, and the relative amounts of each are adjusted for a compromise between sluggish action and stability.
- Distance-velocity lag
Delay of a signal for a period of time will give a graph of the inlet that is exactly the same as a graph of the outlet except for a displacement in time. For example, plug flow of a solution through a pipe may have negligible mixing or dispersion, thus an element of fluid may be assumed to traverse the pipe unchanged. Distance-velocity lag is a killer in terms of process control. Consider a sensor in a flow system where the sample takes time to travel from the bioprocess to the sensor. If the process variable is increasing and decreasing, the signal from the sensor can be out of phase so that the control action can be backwards from the actual process condition. The action that was supposed to make the process better can be in the wrong direction and make it worse. The way to reduce distance-velocity lag is to put sensors right in the vessel, to reduce distances between units, or to pump more rapidly through the connecting lines.
- Feedforward control
The basic idea is to estimate the future error by calculating from the signals now. Feedforward control requires a model of the process. The model need not be exact because updated information from the process can correct for deficiencies. The better the model, the further into the future it can predict accurately.
Practical controllers often mix feedforward control with feedback control. This can compensate for actual errors while allowing for estimated future errors. The models for feedforward control can be elegant systems of simultaneous differential equations or rather simple rules of thumb. Linguistic models supplemented by rules based on artificial intelligence can be quite powerful.
- Forcing functions
The mathematics of process dynamics are derived from idealized signals. A *step* input changes to a new value suddenly. A step that returns to its starting value quickly is called a *pulse*. If the pulse has very short duration, it is an *impulse*. A *ramp* input changes linearly with time and is shown as a line sloping either upward or downward on a graph. A typical periodic input is a sine wave moving above and below its average value. Inputs to real processes may be approximated well by these idealized functions, or real inputs can be transformed by such methods as Fourier Series Analysis to a combination of sinusoidal inputs.
- Integral control
The error (the difference between the actual and desired condition) is integrated and determines the amount of corrective action. Corrective action

thus accumulates to drive the offset to zero. Seldom is purely integral control used; a combination is common by which proportional control dominates and integral control minimizes the offset.

– Proportional control

Corrective action is in direct proportion to the error. For example, if the bioprocess were far below the desired temperature, the heat would be turned to full on. As the system approached the set point, heat input would be decreased. Because there must be an error to get any corrective action, proportional control has a fundamental offset; there will always be a small error. Although the error can be decreased almost to zero by employing large gain (the multiplier in the transfer function), this may cause instability. Steady-state gain is the multiplier when the transients have died out.

– Set point

The value selected as the target for control. The simplest controller is an on-off switch that supplies corrective action when a property of a system rises above or drops below its set point. The process response usually overshoots and oscillates above and below the set point. For sensitive processes, this variation may be unacceptable, and better control modes must be employed.

– Transfer function

The response of a system to an input function depends upon the governing equations, and the expression that is used to derive the output from the input is called the *transfer function*.

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Automation of Industrial Bioprocesses

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The dramatic development of new electronic devices within the last 25 years has had a substantial influence on the control and automation of industrial bioprocesses. Within this short period of time the method of controlling industrial bioprocesses has changed completely. In this paper, the authors will use a practical approach focusing on the industrial applications of automation systems. From the early attempts to use computers for the automation of biotechnological processes up to the modern process automation systems some milestones are highlighted. Special attention is given to the influence of Standards and Guidelines on the development of automation systems.

Keywords. Automation, Biotechnology, Process control, Computer control, Computer validation.

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1

Introduction

1.1

Characteristics of Bioprocesses

In the context of this study, the spectrum of bioprocesses is restricted to transformations of substances by microorganisms or cells in submersed cultures on an industrial scale to achieve one or more of the following goals: 1) degradation of complex substances into simple components, 2) synthesis of substances which may be accumulated in the microorganisms or excreted to the medium, 3) production of biomass from some nutrients. Usually the processes run in some kind of bioreactor to guarantee more or less homogeneous conditions and to perform the mass transfer of gaseous components creating the necessary turbulence.

Depending on the characteristics of the process and on the set goals, the process is carried out under sterile or non sterile conditions as a batch or continuous cultivation of one or more strains, the latter running in a stationary or, in some limits, in a non-stationary state. The different types of bioprocesses have different control requirements. A large-scale continuous culture for yeast production running in a stationary state is easily controlled by some parameters, such as temperature, pH, aeration and dilution rate; there is no demand for any logistics. In biological wastewater treatment running under continuous, non-stationary conditions, decisions are made depending on the highly variable input substances or on various maintenance requirements. Some type of logical reasoning has to be introduced into process control.

The two examples mentioned above would not justify a distinction between the control of a bioprocess and any other chemical process. However, regarding typical biological batch processes in the field of pharmaceutical production, running under rigorous sterile conditions, e.g., the production of insulin, interferon or vaccines, a high degree of complexity and special requirements justify a distinct discipline to describe the control of biotechnological processes. These processes consist normally of 5 main phases:

- equipment check,
- sterilization,
- cultivation,
- downstream,
- cleaning.

The plant itself is composed of dozens of piped vessels, hundreds of control loops, thousands of valves and many additional devices. The process is hardly in a stationary state and the setting of the entire periphery has to follow a recipe

for the specific product. Exception states and alarm situations have to be mastered to protect human lives and costly equipment. Additionally, the validation of pharmaceutical processes and legal prescriptions requires sophisticated documentation describing all the details of each batch. In a modern biotechnological plant such processes run automatically.

1.2

Nature of Processes Automation

Today, automated devices are found in everyday life. From the standpoint of common sense, men are replaced by machines in automated systems. In earlier times automation was based on mechanics, as for example an electric piano, a machine fed by a program punched on paper rolls replaced the pianist. Rows of holes are scanned sequentially producing the time axis and depending on the row position an actuator was activated. Every activation turned the playing on or off. This simple example shows the basic characteristics of an automatic process: The system consists of an information processing machine communicating with a user (start/stop knobs), a process interface (driver of the piano hammers) and a program (punched paper rolls) evaluated stepwise. Depending on the automatic process the information processing is uni- or bi-directional. Therefore, the automated process can take into account process data inputs or establish a dialog with the user.

The basic element in automation is *the control loop* used to adjust an actual value to a given setpoint automatically. The feedback mechanism introduced by control loops is of primary importance for technical processes and, in industrial process control, many of these basic elements are needed. Like any automation, the control loop consists of a set of inputs, outputs and a program, however certainly on a level of lower complexity compared to the control of a whole process. The possibility to structure process control into different hierarchical levels is of primary importance, as will be explained in the next section.

1.3

Structural Design

Regarding the previous examples of pharmaceutical production, processes are composed of thousand of devices, the implementation of automation is only possible on the basis of a well-established structure using a system of one or more computers including process and user interfaces, program libraries and data bases. The list of process interfaces defines all of the I/O-channels available between the process control system and process field. The operations that can be accomplished have to be functionally described at least in three hierarchical levels:

Control Level 1. At the lowest level, I/O-Channels are grouped together to control loops, as for example the control of temperature, pressure, gas and fluid flow or pH-value. Control is realized by looping step-function sequences representing in most cases a PID-algorithm with a constant scanning interval. Therefore the

computer has to guarantee a deterministic time behavior otherwise the PID-algorithms will not work correctly. The control loops can be switched on and off by the operator using the user interface. If the user interface allows the setting of all the outputs and displays all the inputs, the process could be run “by hand” switching on and off the respective control loops or output devices and adjusting the setpoints to the required values. This level is called DDC-level (Direct Digital Control).

Control Level 2. The next higher level defines logistic autonomous process units with the capability of performing a set of operations. Process units may be represented in a hardware structure, like bioreactors, transfer pipes, medium tanks, downstream equipment etc., or they can represent an abstract logical unit, like a scheduler servicing CIP-requirements of other process units. The same operation can run with different sets of parameters. By selecting the right parameter-set, the operation can be adapted to a specific product. An operation may start only under some defined conditions. Locking mechanisms are based on the state of the process unit, and exception routines are required to avoid disastrous situations. According to the functional description the operation has to document its own execution. At level 2, the requirement concerning the time behavior is much less critical compared to the DDC-level, access to mass storage devices and network communications should, however, not disturb the operation’s execution. With a combination of level 1 and 2, processes of medium complexity can be automated. The user starts the needed operations, in the right sequence following a written recipe or Standard Operation Procedure (SOP). For a continuous process typically only a few operations have to be started. However in a batch process, the user has to control a large number of sequential operations, starting each operation and waiting for its completion. As this is very time-consuming, manual procedures are not satisfactory for complex production processes and a third level of control and automation is needed.

Control Level 3. The third level represents a scheduler starting operations according to a recipe. Formally, the recipe may be represented by a graph, where each node corresponds to an operation waiting for its start after the termination or completion of one or more running operations. If an operation fails, the corresponding process unit will not reach the end state that the scheduler is waiting for. A simple strategy to handle such exception states is based on a manual restart of the failed operation. As soon the corresponding process unit reaches the correct end state the scheduler will continue and start the next operation.

This short description of the three levels illustrates that by proper structuring the automation of complex bioprocess can be managed. There are still, however, many more important problems left to consider. Some of these are universal in computer applications, such as the problem of creating an optimal “Human-Machine-Interface”.

It is obvious that a successful implementation of a control system depends on an adequate documentation comprised of at least the user requirements, functional descriptions, program code, and test procedures. This implies that a

rigorous quality management system should accompany the implementation as it is now defined for the pharmaceutical industry in the Suppliers-Guide from the GAMP-Forum [1]. Following these guidelines a firm base for validation is achieved.

1.4

Problems of Hardware Architecture

The structural principles outlined above can be realized with many different hardware architectures, computers, and process periphery. The architectures differ mainly in the degree of centralization.

The process interfaces can be connected directly to the computer and the wiring to the transmitters may be realized in the form of a star topology. On the other hand, a decentralized solution could consist of a fieldbus system with interfaces distributed throughout the whole process area and connected serially to the fieldbus. In a typical fieldbus controller; the bus topology is mirrored in the controller's memory and contains the actual value of each device updated in time intervals of some milliseconds. The computer no longer has to directly access the interfaces. Today, fieldbus systems are widely accepted: the cabling is transparent and allows for easy maintenance, the interfaces are close to the equipment and decoupled from the computer, the computer itself has only to read from and write into memory cells inside the fieldbus controller to exchange data with the process periphery. Such systems are extremely highly reliable and have sophisticated error-detecting facilities built in.

The question concerning the "right" computer system architecture is discussed among suppliers and users in a controversial manner. In the past, process computers depended on low performance processors with low storage capacities (memory, disk devices). Consequently the development of process control systems led to distributed systems with a lot of small computers controlled by one or more master computers. However the performance and capacity of today's computer systems and the availability of powerful real-time operating systems allow the realization of process control systems of any degree of complexity in one single computer. For today's pharmaceutical applications, reliability is probably the most important requirement. But it is very difficult to compare the reliability of a multi-component distributed system requiring a considerable amount of network traffic to a single component system without a network to other process computers. Maybe nature itself has given the answer to this problem. During their evolution biological systems ended with a centralized computer system, and a decentralized peripheral system. Perhaps a process control system should evolve in the same manner. Moreover a centralized computer system consisting of two redundant computers and a redundant fieldbus system seems to be an optimum solution. But as the main requirement for industrial applications, the systems should be as simple and robust as possible. Furthermore many interesting developments have been developed in the academic world (sophisticated controllers, process optimization, application of Artificial Intelligence (AI) etc.) are still too sensitive to survive in the daily life of an industrial production process.

1.5

Benefits of Bioprocesses Automation

- Automation makes it possible to run bioprocesses of any degree of complexity.
- The recipe-controlled batch guarantees a product of constant quality accompanied by the necessary documentation of the production process. At any time, the batch corresponding to an individual product can be traced back to its origin. Accurate documentation is needed to fulfill legal or validation requirements.
- Automation increases the reliability because the operator is supported by the automation system (check lists, alarm messages, help libraries).
- The safety for humans and materials during the production process is substantially increased because an automation system checks the critical parameters continuously. In addition the system is capable of handling failures according to defined exception routines.
- The economics are improved as time and personnel are saved.

2

History of Automation of Industrial Bioprocesses

2.1

General Considerations

The term automation may be ambiguous. The meaning may be automatic control (e.g. control of temperature), but automation, as it is understood by the authors is control on a higher level and involves sequences of tasks based on a schedule or a series of events as defined by Singh et al. 1990 [2]. In this historic retrospect, we will mainly focus on automatic operation.

The history of the implementation of automatic operations into biotechnological processes may be as old as the history of biotechnology. It was probably always man's intention to replace repetitive operations by machines. However modern automation developed parallel to the development of the technologies and equipment needed for process automation. It may be generally observed that in industrial biotechnological processes, technologies and equipment available on the market have been adapted to the special requirements and implemented. There are only a few exceptions where control and automation equipment were specially designed and developed for biotechnological processes. It is only the requirement for sterile operation of biotechnological processes that asked for special developments. In Biotechnology, sensors and actuators have to be able to withstand sterilization conditions and as sensors and actuators are an essential part of any automation system a few words on the history of these components will be given in the first part of this historical consideration.

In theory, process automation is only possible if the process itself is known and the process behavior can be predicted at any time. Therefore, besides the availability of control and automation equipment, it is an important necessity

that these two requirements be thoroughly understood and respected. For biotechnological processes these may still be the weakest points in the attempt to automate bioprocesses. A few considerations to these problems will be mentioned in part two of this retrospective.

Automation needs equipment capable of acting according to preset patterns or algorithms but there are also logical components needed to automate a process. The development of logical devices from simple relays to modern process computers had a continuously strong influence on the development of automation systems. In Sect. 3, this influence of new electronic control equipment on process automation in biotechnology will be reviewed.

To summarize, successful automation of industrial bioprocesses is only possible if 1. the equipment (sensors and actuators) for sterile processes is available, 2. if the process in its basic behavior is known and predictable and 3. if the controllers with the needed algorithms are available. An attempt to illustrate the history of bioprocess automation has to take these points into consideration. The following are prerequisites for bioprocess automation:

- Availability of Field equipment suitable for sterile operation (sensors and actuators);
- Known and predictable Process Behavior;
- Availability of reliable Automation Systems (Computer Systems and Software).

2.2

Process Control Equipment for Sterile Conditions

Even in the first industrial bioprocesses for the production of antibiotics in the late 1940s, automatic features such as control loops for temperature and impeller speed had been implemented. As at this early stage of industrial biotechnology, sterility was not absolutely necessary, techniques developed for other applications could easily be adapted to bioprocesses. However, with the requirement for sterile conditions the need for special sensors and actuators arose. As an example, the measurement and control of the pH-value should be mentioned. In this context, the development of the first sterilizable pH-electrode by Fiechter et al. 1964 [3] has to be considered as an important milestone in the history of bioprocess automation. As a result of this pioneering work, one of the most important parameters for biological reactions could now be measured and controlled under sterile conditions. Of comparable importance on the way to an automated bioprocess was the development of membrane valves capable of operating under sterile conditions as introduced by several equipment manufacturers in the 1970s. In the scientific literature, these valves have not been considered worth mentioning. However, only these valves allow for an interaction with the process while maintaining sterility and it is hard to imagine any modern biotechnological processes without them.

These two examples may be representative of the importance of suitable field equipment for the automation of bioprocesses. They are mentioned to illustrate that process automation comprises not only electronics and computers. Without sensors and actuators, even the most sophisticated computer system would be useless.

2.3

Known and Predictable Process Behavior

In theory, automation is only possible if the process behavior is known and predictable at any time. Although knowledge about biological reactions has increased immensely during the last decades, it never would allow us to interpret the extremely complex behavior of biological systems. With its huge variability, a biological process is not predictable. The on-line measurements do not contribute much to overcome this lack of knowledge as there are still only a few exceptions known where biological quantities such as biomass, products, intermediates or substrate can be measured on-line in an industrial environment. Consequently far more than 90% of the scientific publications about "Bioprocess Control" focus on these problems. New analytical procedures, new sensors, process and control models, optimization and its implementation into control strategies, dominate the scientific literature. As soon as the first mini-computers appeared on the market, biotechnologists all over the world used computers to calculate process parameters based on various process models. With the appearance of personal computers this tendency even increased. The recently published Proceedings of the 7th IFAC International Conference held in Osaka from 31st May to 4th June, 1998 [4] gives an excellent overview on the present status of research and development activities in this field. It is not the authors' intention to review these scientific research and development activities. These are certainly of great importance for the understanding of biological reaction systems and they may be used once in future automation systems. Up to now, none of these models have been implemented into real industrial production processes. Industrial biotechnology seems to operate pragmatically and does not care about the lack of basic knowledge about biological reaction systems. By dividing the whole process into smaller process units with known behavior, by taking off-line data and experiences into the control concept and with a combination of automation with manual interactions, a high degree of automation can be achieved despite the fact that the detailed behavior of the process itself is not known. This proves that the limiting factors for successful automation of biotechnological processes are more technical rather than biological.

2.4

Use of Automation Systems in Industrial Bioprocesses

In addition to the previously mentioned prerequisite of the availability of field instrumentation and process knowledge, successful process automation requires a logical device to control the operation according to a preset. Even in the very first industrial bioprocesses for antibiotic production, the technical equipment to control and to automate these processes was available. Analog controllers mostly configured for PID-Operation (Proportional-Integral-Derivative) had been used to solve all of the control tasks. Logical elements such as timers and relays had been allowed to fulfill the automation requirements. As these hard-wired techniques were very difficult to realize and to maintain, automatic

operation was normally limited to defined small standard operations such as sterilization, product harvest or media transfer. Automation of complete production plants or even recipe handling was not feasible with this hard-wired techniques.

This changed completely when, in late 1960, a newly designed solid-state controller was introduced into the process control market. This new device, called a programmable logic controller (PLC), not only replaced the relay logic controllers, but more importantly offered new functionality not yet realized with conventional analog controllers. These PLCs were quickly implemented into biotechnological plants; at the beginning just replacing the conventional relay logic. All leading plant manufacturers at that time realized standard operations with PLCs.

The PLC is functionally divided into four parts: the input, the output, the logic unit and the memory unit. This basic principle has remained valid until now although the PLC has become much more powerful (more memory, speed) and flexible (more functionality) in the last decades. Still, PLCs are widespread in biotechnological plants and are used to do much more than simple control sequential actions. Whereas single stand-alone equipment (such as a centrifuge or a filtration unit) is relatively simple to automate with a PLC, the automation of complete plants comprising several bioreactors, tanks, up- and downstream equipment is not within the PLCs reach. The immense effort to coordinate the actions of single PLCs to handle a recipe that requires multiple devices and various equipment may end in an immense traffic jam in communication. In the extreme, each PLC has to mirror the status of all the other PLCs in the same production unit. Consequently, today process computers are replacing PLCs more and more.

In the 1960s, the general-purpose digital computer was brought to the market and soon after also applied to biotechnological applications. In contrast to PLCs, these general-purpose computers offered a complete versatility irrespective of the application. The functionality was defined by software alone. Additional features such as mass storage, communication networks, visualization devices as well as an operating system controlling the interactions of the different modules, were now available. The initial differences, mainly based on performance and price, between micro-, mini- and mainframe computers has decreased more and more over the last two decades. Biotechnological companies and research institutions recognized very quickly the great potential of these universal computers and used them to acquire and to store data, to control process parameters, and to automate operation sequences. Furthermore due to the high calculating power of these machines, on-line process modeling became possible. A favorite among the computers used in those early days was the computer series PDP 8 to 11 (Programmed Data Processor from Digital Equipment Corporation).

No publications have been found showing the industrial use of these mini-computers in the early days. Therefore the authors contacted all leading biotechnological companies as well as manufacturers of biotechnological equipment to get information on the early use of computers for process automation. Unfortunately only three answers have been received from more than

50 requests sent out. On a follow-up by phone, the authors received mainly the same answer: all relevant data had disappeared. It seems that during these dynamic developments of the last three decades, nobody considered the historical value of data and equipment. Therefore, this review is mainly based on the personal experiences of the authors and may not correctly represent the whole situation.

New Brunswick Scientific Co. Inc., probably the first plant manufacturer offering bioprocess systems controlled by minicomputers, provided us with some pictures of the early days of computer applications in biotechnology and we feel it worthy enough to publish these pictures as historical documents (Fig. 1). Looking at these pictures it is hard to believe that there are only 25 years of development between the system shown and a modern process automation system used today. The computer system to control a relatively simple pilot plant needed a complete room, which also had to be air-conditioned. Memory at that time was limited and the programmers were forced to optimize their code in order to save space. A very efficient real-time operating system organized the available memory of 128 kbytes in a way that much larger applications could be executed successfully. The “Human-Machine-Interface” was at that time a video terminal with a keyboard and the information displayed was completely text based. No on-line graphics display was available at that time.

This period characterized by the use of the PDP11, may be considered as a real milestone not only for the development of automation systems for biotechnological applications, but also for the general understanding and further development of the whole biotechnological industry. It initiated a remarkable change in the mostly biology oriented biotechnology of that time. From that time on, natural science was definitively influenced more and more by engineering sciences and biologists had to learn to communicate with engineers. Biologists had been forced to describe the “Art of Fermentation” and to convert their experiences into Bits and Bytes. The way of looking at a bioprocess had completely changed.

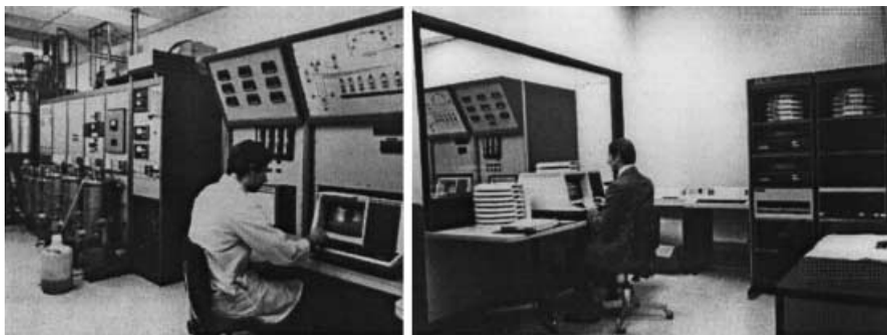


Fig. 1. Control of a biotechnological pilot plant in 1978 by a PDP11 computer system. With courtesy of New Brunswick Scientific Co. Inc.

The period of the PDP11 was only of short duration. New computer generations arose with much more powerful components (e.g. HP-1000 series from Hewlett Packard, Honeywell 4500 or DEC Vax Series) were soon brought onto market. Fortunately, this development of new computers went in parallel with a generally prosperous growth of the pharmaceutical companies. Some of these companies started strong investments into biotechnology and during the late 1970s and early 1980s many new pilot plants and production facilities were built. All of these plants had been equipped with the latest computer products available at that time. Reports have been published on installations at Elli Lilly and Company [5], Genetic-Institute [6], Merck Sharp and Dohme [7, 8], Schering-Plough [2] and Smith Kline [9]. All of these biotech companies elaborated on the use of computers to control and to automate their bioprocesses. They demonstrated not only in different ways the principle functioning of computerized biotechnological plants, but added substantial new features to the systems. The focus of that time was no longer on just the control and automation but also on the evaluation and management of process data. Process computers were networked, supervisory systems implemented and communication with company information systems was realized. The networked bioprocess was born. Simple graphic displays started to replace the text based "Human-Machine-Interface" and made the system accessible to people who were not computer engineers.

Common to all these installations was a very high investment in equipment and manpower needed to keep the system running and it is obvious that this restricted the numbers of possible users. With the introduction of the Personal Computer (PC), in 1977 by Apple, computers were made more affordable and available to a broader range of companies and institutions. The age of PC-controlled processes was born. In research institutes all over the world, activities went on to couple bioreactors with PCs. Pioneering works on the use of PCs for on-line calculations of biological process parameters were published by Röhr et al. 1978 [10]. One of the first PC-based bioprocess systems commercially available on the market was developed by B. Braun Biotech International GmbH, Melsungen [11] 1982. Based on an HP-85 desk-top PC, set-point-control (SPC), data acquisition and visualization, as well as automatic sterilization and even some simple on-line calculations of biological properties such as RQ-value had been realized. It is not known to the authors how many of these systems were sold. It may have been only a few units because the technical improvement of PCs was so fast that shortly after the appearance of a new system on the market, the products themselves were already obsolete. However, historically this example may be of considerable importance as it stood at the beginning of a new trend for equipment manufacturers. Since that time, all of them have had to develop and maintain PC-based control systems. Still today in smaller installations PCs are used to control and automate bioprocesses. Together with the very powerful control and automation software available now, the PC has developed into a wide spread automation device and has replaced more and more the traditionally applied PLCs.

Attempts to automate complete industrial biotechnological production plants have only been undertaken since the late 1970s and were initiated by the

Table 1. Functional levels of process automation systems

Functional level	Main functions
Data Communication Highway (ETHERNET)	Communication with further information systems
Supervisory computers	Plant automation, recipes handling, data bases
Operator Interfaces	Operator interactions with the process
Front-end Process Computer	Real-time control and automation of single process units
Sensor/Actuator Fieldbus	Field data handling communication with process computer
Process	Process 1 to n with Sensors and Actuators

powerful process computers which appeared on the market. In 1975, Honeywell introduced the first process computer system (TDC2000) and shortly after this system was used in a biotechnology plant. With this system a standard was set. All leading automation companies such as Siemens, ABB, Fisher-Rosemount, Foxboro, just to mention a few, soon brought comparable systems onto the market.

In parallel with the development of new process computers, different companies elaborated new concepts for field instrumentation. Instead of the traditional centralized communication of sensors and actuators with the computer system, fieldbus systems for decentralized communication with sensors and actuators were developed. The biotechnological industry soon realized the great advantages of this fieldbus concept and implemented it in plant automation systems. Whereas in research and laboratory environments, the CAN-Bus is often used [12], in industrial biotechnological plants, mainly Profibus and Interbus-S can be found.

The general concept of a modern up to date automation system for biotechnological processes comprises many of the different functional levels shown in Table 1.

Depending on the computer hardware applied, different functional levels may be implemented into one computer system.

2.5

Some Aspects on the Development of Programming Languages

For the very first computer based systems, the programs had to be written by the users themselves. The programming language at that time was mostly FORTRAN. Additionally, some time-critical tasks had to be written in Assembler (a CPU-architecture-dependent low-level language). Although there are still some FORTRAN programs around, FORTRAN was replaced more and more by languages like Pascal, C and later C++. Together with the PCs, the BASIC language was developed and most PC-based applications had been written in this language. It soon turned out that the "Spaghetti-Code" of BASIC programs was not appropriate for complex applications. BASIC disappeared

almost completely and is replaced by either Visual Basic or commercially available control and automation packages today.

Writing the application software was the crucial point in all automation projects and there were practically no projects where the costs and time for software development had not been underestimated. At the beginning of the computerized age, the industrial users set up their own automation groups and developed the software in-house with the important disadvantage that all the efforts made could normally only be applied once. This may be the reason that more and more specialized automation engineering companies emerged. These companies first started with their own software developments but more and more went over to base their application on generally available control and automation software packages.

Furthermore, equipment and plant manufacturers started to provide their equipment with their own automation software packages such as B. Braun Biotech International GmbH, Melsungen does with the MFCS for bioreactor automation [13] or New Brunswick Scientific Co with the AFS-BioCommand. These Programs may serve quite well to automate the equipment of these companies but can hardly be applied to other manufacturers equipment or to automate complete plants.

The appearance of universal configurable software packages for process control such as Genesis, Lab-View and others may help to minimize investment in time and money. Examples of the successful use of such commercially available automation packages have been published by different authors [14, 15]. All of these packages offer graphically oriented configuration tools and are relatively easy to handle, also by non-specialists. In addition, such software packages offer the advantage that the user normally does not have to care about future upgrades and compatibility with new computer or operating systems. It may be assumed that the software company takes care of this. A disadvantage may be the dependency on the software supplier. If a special need of the user is not part of the package it is nearly impossible to fulfill this need.

Large automation companies offer software solutions together with their computer hardware products. All of these general automation software packages can be used for biotechnological applications. However, the configuration of the software needs automation specialists to convert the functional process description into program code. Very intense communication between the biotechnologically oriented user and the general automation engineer is essential for a successful realization. There are projects known to the authors where famous automation companies failed to automate biotechnological processes, not because of a general hardware deficiency but as a consequence of a communication problem between the biologically oriented users and the technologically oriented automation engineers.

Although the history of automation of industrial biotechnology is only about 25 years old, it is extremely difficult to structure and to weight the data. The history does not comprise single events that can be chronologically ordered but consists of complex interactions of various parameters (Table 2). The development of new computer products was certainly one of the driving forces, but it was not the only one. Of similar importance may be the biotechnological in-

Table 2. The evolution of computers and process automation systems for industrial biotechnological application

Computer Generation	1. and 2. Generation Experimental and Central Batch Computers	3. Generation Centralized Computers with Direct Dialog Interactions	4. Generation Personal Computers and Workstations	5. Generation Parallel Computers Client Server Systems Multimedia Computers	
Computing Performance MFlops					
Storage Capacity	Kbytes	Mbytes	GBytes		
Operator Interfaces	Punch Cards	Text Display on Video Terminals	Graphic Display on Video Terminals		
		Key Board	Mouse, Barcode Reader		
		Line Printers	Graphic Printers, Color Printers		
Type of Automation System					
Typical Programming Languages	Assembler				
	Fortran, C, Pascal				
	BASIC				
	Object oriented Languages: C++, ObjectPascal ...				
	Logical and Visual Languages				
Type of Application	DDC (Direct Digital Control)				
	SCADA (Set-Point Control and Data Acquisition)				
	Batch Control				
Year	1960	1970	1980	1990	2000

dustry itself with its special requirements for process automation providing pressure for new system developments. The extremely dynamic computer market, the rapid appearance of new software developments and last but not least the dynamics of the biotechnological industry itself, makes it difficult to draw conclusions for future tendencies. The trend for future computer developments will still go on: *smaller, faster and more powerful!* On the other hand there is the industrial need for *reliability and continuity*. This will slow down the integration of new hardware into process automation. In addition the increasingly stricter regulations and *standards* for automation systems for the pharmaceutical industry will greatly influence future developments. It may therefore be concluded that the generation time for industrial automation system will still be much larger than the generation time for computer hardware.

3 Standards and Guidelines

3.1 Why Standards?

Before proceeding into details of the different standards involved in process control automation, there is a brief discussion about standards themselves. Why are standards needed? Who determines that they are standards? Who enforces compliance with standards?

Standards are necessary in order to engage in commerce of any type. Component parts for any kinds of “final product” are manufactured at many different locations by different vendors. Each individual contributor involved must design, manufacture and test his components according to the specifications he received. In the “final product”, assemblies of different parts must fit and work together as a whole as specified by the customer, even though the individual contributors may never have even met. Without common agreement, common reference and common understanding, such a procedure would not be possible. Therefore, national standards have been developed in all major industrial nations. The units of the standards might differ from nation to nation; however, conversions between them are possible as long as everyone concerned understands the standards themselves and as long as those standards selected remain constant. The need for international standards has been recognized by different national standard organizations and has given birth to international non-governmental bodies such as the International Electrotechnical Commission (IEC) founded in 1906 and the International Organization for Standardization (ISO) established as an organization in 1947. National or international vendor or user-groups set up other standards. To go into detail concerning the specifics of the different organizations and standards is beyond the scope of this paper. However, some standards related to the process control systems will be detailed further in the next sections.

3.2 Computer Directives History

Regulatory authorities around the world have come to recognize the importance of electronic data systems used in the research, development and manufacture of products and services coming under their jurisdiction. For more than a decade, authorities have been issuing guidance documents for the industries and to assist their inspectors in how to review systems for Good Practice compliance. The percentage of computer content in regulated activities has been growing rapidly since 1980. With this growth in the use of computers has come a parallel growth in regulatory guidance for computer validation to assure the quality of electronic data handling and electronic process controls.

With the growth in regulatory directives, forums were founded to promote the understanding of the regulation and use of computer and control systems

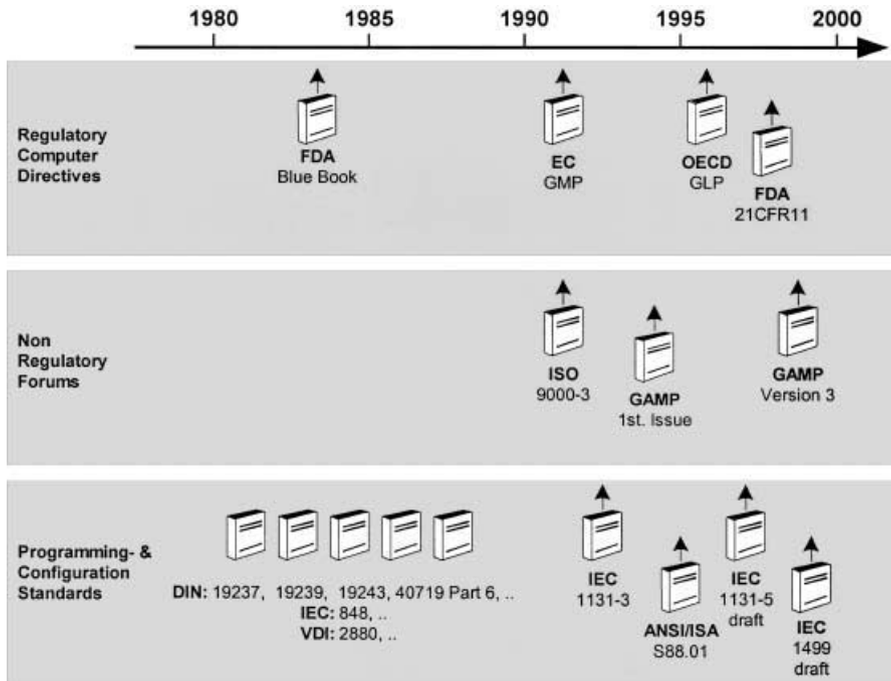


Fig. 2. History of computer directives and standards

within the pharmaceutical manufacturing industry. Different “non-regulatory” standards were also established by such forums.

Independent of regulatory issues, the industries and the national- and international committees agreed to standards defining programming and configuration issues in computer control. Figure 2 illustrates the history of process control computer directives and standards. These “documentary milestones” detailed further in the following chapters represent the personal point of view of the authors, as the result of more than twenty years of involvement in the field of automation of biotechnological applications.

3.3

Regulatory Directives

1983: The Blue Book. The first document published in 1983 became known by the color of its cover – the Blue Book [16]. Its focus is on computerized systems used in drug manufacturing and instructs inspectors of the US Food and Drug Administration in computer characteristics and how to inspect systems under the regulations for Good Manufacturing Practices (GMP). The purpose of the guide is to provide the field investigator with a framework upon which to build an inspection of drug establishments which utilize computer systems. The document was not intended to spell out how to conduct a CGMP drug in-

spection or set forth reporting requirements, but rather what aspects of computerized systems to address during such inspections and suggestions on how to address the systems. The guide discusses some potential problem areas in application of computer systems, provides inspectional guidance, and includes a glossary of terms the investigator should be aware of prior to performing the inspection.

The blue book triggered mystification among the manufacturers and the suppliers of automation systems. FDA validation advanced to a key issue in each automation project, even in Europe. The fact was that most people had problems in transforming this guidance into their own concise quality and validation plan. The subsequent issue of supplemental compliance policy guidelines [17–23] and the Pharmaceutical Manufacturers Association (PMA) Computer System Validation Committee (CSV) concept paper [24] are the basis for today's computer system validation practice.

1991: GMP – Europe. The European Community adopted directives laying down principles and guidelines of good manufacturing practice (GMP) for medicinal products. The Guide to Good Manufacturing Practice [1] is used to assess applications for manufacturing authorizations and is a basis for inspection of manufacturers of medicinal products. The first edition of the guide was published in 1989, including an annex on the manufacture of sterile medicinal products. The second edition was published in January 1992; including the Commission Directives 91/356 of 13th June 1991 [25] and 91/412 of 23rd July 1991 [26] laying down the principles and guidelines on good manufacturing practice for medicinal products for human use as well as for veterinary medicinal products. The second edition also included 12 additional annexes. The basic requirements in the main guide have not been modified, but 14 annexes on the manufacture of medicinal products were included in the 1998 edition.

Annex 11 [27] to this guide gives a concise view of good computer practice in nineteen paragraphs of directives covering the training and experience of personnel, the extent of validation throughout the whole life cycle of a system, and operational guidelines for system development, deployment, risk management and supplier contracts. Annex 11 reflects four major themes:

1. *Management Control*: Does the company apply System Life Cycle activities?
2. *Data Integrity*: Is electronic data handled properly?
3. *Reliability*: Is the systems continuous performance reliable?
4. *Auditability*: Is there documented evidence for all above topics?

1995: OECD GLP Consensus – Europe, Japan & US. The Organization for Economic Development and Cooperation (OECD), which includes Europe, Japan and US published a consensus document for applying GLP (Good Laboratory Practice) principles to computerized systems [28]. The OECD document contains more as Annex 11, and also adds other items of concern such as the definition of raw data and retrospective evaluation. A more detailed comparison of validation guidance has been issued in a white paper by the Digital Equipment Corporation (DEC) [29].

1997: 21 CFR 11. In 1997 the FDA issued regulations that provide criteria for acceptance by the FDA, under certain circumstances, of electronic records, electronic signatures, and hand-written signatures executed to electronic records as equivalent to paper records and hand-written signatures executed on paper [30]. These regulations, which apply to all FDA program areas, are intended to permit the widest possible use of electronic technology, compatible with FDA's responsibility to promote and protect public health.

3.4

Non-regulatory Forums

Several groups around the world, including the Instrument Society of America (ISA), the International Electrotechnical Commission (IEC), the International Organization for Standardization (ISO), the GAMP group, Germany's Normenarbeitsgemeinschaft Mess- und Regelungstechnik der Chemischen Industrie (NAMUR) and others are working on improving and standardizing different fields of process automation.

1991: ISO 9000–3. ISO prepared the 9000–3 guideline [31] in order to help organizations to apply the ISO 9001 standard to computer software development, supply, installation and maintenance. The ISO 9000–3:1997 edition is actually an expanded version of the old ISO 9001:1994 standard. ISO has simply copied the old text from ISO 9001 [32] and pasted it into the new version of ISO 9000–3, and then added some new text that refers only to software.

The spirit of ISO 9000 is that customer and suppliers should work together to guarantee consistent quality standards and to have procedures in place so that when the quality standards move away from those the customer requires, corrections are made. However, ISO 9000 allows the supplier to have procedures in place, without really taking into account the customer's needs. ISO 9000 is an accreditation system, which lacks technical depth.

1994: The GAMP guide. With the increased penetration into, and complexity of automated systems in pharmaceutical manufacturing, the focus on such systems has increased. With it comes the need to improve the understanding of the regulations and their interpretation. Better communication was required, not only within the pharmaceutical industry, but also with its suppliers. Thus an informal group, the UK Pharmaceutical Industry Computer Systems Validation Forum (now known as the GAMP Forum) was set up to promote that understanding. Draft guidelines entitled "Good Automated Manufacturing Practice (GAMP)" were created by the forum led by this forum. Its remit was:

1. To devise a draft set of guidelines for suppliers of automated systems to the pharmaceutical manufacturing industry.
2. To take account of the requirements of both the European and North American regulatory bodies.
3. To make use of existing internationally recognized standards where appropriate.
4. To consult with the Medicines Control Agency in the UK.

The first draft was approved by the Forum and made available for comment from the suppliers of automated systems to the industry and other interested parties in 1994. The current version, GAMP 3 [1], enables compliance with the requirements of the European Community as explained in the European Guide to GMP including Annex 11 Computerized Systems”, and in particular item 2 (validation and the Life-cycle) and item 5 (software quality assurance) for system development and validation. Systems designed and implemented in accordance with this guide should also meet all the other requirements of the EC guide. This approach should also be acceptable to the FDA.

The Guide is built around a formal management system for the development, supply and maintenance of automated systems by a supplier. Adherence to this management system by the supplier will provide both the system and sufficient documentary records to enable the complete system to be accepted and validated by the user. The formal acceptance of an automated system and its documentation by the user is an integral part of the validation of that system. This formal acceptance by the user is a key issue. Due to the fact, that this documentary evidence is required by the pharmaceutical companies as an integral part of validation, it is defined as: “Establishing documented evidence which provides a high degree of assurance that a specific process will consistently produce a product meeting its pre-determined specifications and quality attributes”.

The actual GAMP 3 guide is like a fill-in-the-blanks manual for the supplier and the client and is designed for prospective validation of new computer systems. It helps focus on all the details needed for specification and subsequent verification of a computerized system’s functionality. It also helps to allocate validation support responsibilities between supplier and customer. The guide consists of a Supplier Guide section, a User Guide section and a Best Practice Guide section.

The Supplier Guide section defines a management system model with life cycle and supporting activities for the supplier.

The User Guide section outlines the different user responsibilities and illustrates a possible validation strategy for different types of software products. The Best Practice Guide section details the validation procedures for Information and Control Systems and Systems.

Conclusions. The GAMP guide represents an excellent basis for a modern automation project management. But one question remains: Who will bear the costs? The investments in the development (specification) process are considerable by means of time and money. Tight time schedules and project budgets increase the problem. The consciousness, however, must be present, that the invested money will be paid back over the lifetime of the product.

3.5

Programming & Configuration Standards

1993: IEC 1131–3. Automation system designers are often required to use programmable controllers from various manufacturers in different systems, or

even in the same system. However, the hardware of programmable controllers from different manufacturers may have very little in common. Historically, this has resulted in significant differences in the elements and methods of programming the software as well, to the existing standards and regulations (DIN 19239, DIN 19237, DIN 40719 Part 6 and VDI 2880). This has led to the development of manufacturer-specific programming and debugging tools, which often carry very specialized software for programming, testing and maintaining particular controller “families”.

An international committee was set up to tackle this problem and produced the world-wide standard IEC 1131 [33], the third part of which deals with the PLC programming languages. A major goal of IEC 1131–3 is to remove barriers to the understanding and application of programmable controllers. Thus, IEC 1131–3 introduces numerous facilities to support the advantages of PLCs, even if controllers of different vendors are concerned. Within the standard four programming languages are defined. Syntax and semantics have been defined, leaving no room for dialects. The languages consist of two textual and two graphical versions:

- Textual,
 Instruction List (IL),
 Structured Text (ST).
- Graphical,
 Ladder Diagram (LD),
 Function Block Diagram (FBD).

The Ladder Diagram has its roots in the USA. It is based on the graphical presentation of Relay Ladder Logic. The Instruction List is its European counterpart. As a textual language, it resembles assembler. The Function Block Diagram is very common to the process industry. It expresses the behavior of functions, function blocks and programs as a set of interconnected graphical blocks, like an electronic circuit diagrams. It looks at a system in terms of the flow of signals between processing elements. The Structured Text is a very powerful language with its roots in Ada, Pascal and “C”. It can be used excellently for the definition of complex function blocks, which can be used within any of the other languages. Summarizing, the mayor benefits of IEC 1131 are:

- Standardized languages and standardized programming;
- Increasing software re-usability;
- Ability to use the best language, depending on the application;
- Common platform for understanding control software.

However, the software model and the programming languages described in IEC 1131–3 are mainly applicable on stand alone or loosely coupled PLCs, not on Distributed Control Systems (DCS). For these applications it is necessary to introduce a communication system which is able to handle Distributed Function Blocks. The concept of communication defined in IEC 1131–5 [34] can be used to solve this problem. To exchange data between physically separated devices IEC 1131–5 offers several possibilities, but all of them are quite complicated and without supported synchronization between the function

blocks. To enhance the communication facilities in DCSs by using IEC 1131–3 it is necessary to first extend the software model of the standard.

A definition of function blocks suited for DCS application is proposed by the IEC Technical Committee (TC65), Function Blocks. This IEC 1499 draft [35] defines a generic architecture and presents guidelines for the use of Function Blocks in distributed industrial-process measurement and control systems (IPMCSs). The models given in this Standard are intended to be generic; domain independent and extensible to the definition and use of function blocks in other standards.

1995: SP88. The ISA's SP88 committee first began work on the specification in 1989, and soon S88.01 [36] became the fundamental building block of good communication between software vendors providing – and companies looking to implement automated batch-manufacturing solutions.

Within the manufacturing sector, production processes are typically defined as continuous, discrete or batch. Continuous processes are those in which output occurs in a continuous flow. Discrete processes are those in which output results in finite quantities of parts that can be counted. Batch processes are those in which output yields a finite quantity of material. In batch control, the mixing and reacting of materials are crucial to the integrity of production. Products are determined by a recipe that has a name and contains all requisite information for manufacturing the product. This includes ingredients or raw materials needed, the order of process steps through which ingredients must pass, the conditions of each step in the process, and the equipment to be used in the process.

The SP88 committee began with two fundamental work items: models and terminology, which focused on verbal communication standards for batch solutions; and data structures and language guidelines, which addressed the digital communication issues involved. Work on the first item has been completed, and has become an international standard. Efforts on the second item are still ongoing.

SP88 defines three reference models: *process model*, *physical model* and *procedural control model*. The mapping of process functionality described in the process model with individual equipment provides the possibility to specify unequivocal elements in the procedural control model. The three models allow a systematic breakdown of a process control system into procedures, operations and phases.

The Process Model defines all processes that lead to the production of finite quantities of material by subjecting quantities of input material to a defined order of processing actions using one or more pieces of equipment.

The Physical Model has seven levels, starting at the top with an enterprise, a site and an area. These three levels are defined by business considerations and are not modeled further in this document. The lower levels of the model refer to specific equipment types and are defined by engineering activities.

A Process Cell is a logical grouping of equipment required for the production of one or more batches. A process cell must include all of the components required to run a specific recipe, since recipes cannot cross process cell boundaries.

A *Unit* is a collection of associated control modules and/or equipment modules that carry out one or more major processing activities. Units are presumed to operate on only one batch at a time. Units operate relatively independent of one another. Often in process plants there exist similar sets of equipment and it is useful to describe these sets as “classes.”

An *Equipment Module* is an instance of a specific phase. A phase is a general representation of a group of equipment, which performs a sub-function for a unit. An instance of a phase is referred to as an equipment module. Each phase contains attributes common to all equipment modules of that specific phase type. When a phase is added to a unit it becomes an equipment module. An equipment module maps to the equipment phase in the engineered logic via tag addresses. Only one instance of a specific phase may be added to each unit. When an equipment module is required in multiple units, it may be shared between the units.

The *Procedural Control Model* is defined by means of a hierarchical structure, which can be correlated with the earlier defined process- and physical models. Procedural control directs equipment actions to take place in an ordered sequence. There are four levels to this model:

- The Procedure.
- The Unit Procedure.
- The Operation.
- The Phase.

The utilization of S88.01 modularization brings a degree of efficiency and flexibility that ultimately will reduce the burden on the automation team. Separating the procedural control code from the equipment control code facilitates flexibility. Modularizing the software into reusable phases allows for changes or additions of procedures without ever having to modify and test the existing phase code. A phase is written and validated one time in the life of the equipment. The equipment unit procedure is tested only to ensure that the phases are called in the proper order, greatly simplifying the overall software validation effort.

The related standard SP88.02 (Batch Control Part 2: Data Structures and Language Guidelines), currently in its fourth draft and scheduled for publication next year, describes how data needed for batch control can be stored and exchanged in ways that are independent of the actual hardware or software. It also provides more detail about what a batch recipe actually looks like, an issue that S88.01 does not cover fully. At the moment people tend to use IEC 1131 sequential function charts to describe recipes. IEC 1131 is an excellent standard for programmable controller languages, but was not designed for recipes. S88.02 will attempt to provide a more appropriate method. Plants that are based on batch operations also incorporate some degree of continuous processing, and this too creates problems for S88.01.

4 Outlook

Frost and Sullivan published a new study on the PLC market in January 1995 [37]. Although this study observes the general control and automation market, the relative figures may also be true for the biotechnological industry. According to this study, measured in the number-of-units, the worldwide PLC-market will grow 12.3% from 1993 to the year 2000. The study forecasts that the small PLCs, 30% of the market revenues in 1993, will increase to 35% in 2000, while the revenues for medium and large PLCs will dip from 30% to 26%, over said period. The increased importance of software is also reflected in the figures: software revenues are expected to increase from 9 till 12% over the same period.

Taking into account the rapid developments in hard- and software it is quite difficult to sketch how the next millennium's process control systems will look like. This paper will focus only on two aspects out of a myriad of possible technical topics.

4.1 Integration of Enterprise Resource Planning (ERP) Systems into the Control Level

Integration of enterprise solutions to the control level requires flexibility and openness in tools to solve real world integration problems. The objectives of minimizing special code development making maximum usage of off-the-shelf products and bringing systems rapidly on-line are the goal. One must look beyond the needs of batch execution and select an environment, which allows for quality and maintenance integration.

The integration of ERP into the control systems might include:

- Downloading the production schedule to the control system queues for automatic startup;
- Download specific material reservations;
- Uploading material consumption quantities and lot numbers;
- Uploading semi-finished product quantities and batch numbers;
- Uploading critical product release results that need to follow an order.

In order to further automate the control system integration to handle maintenance and product quality by directly scheduling the control systems, there is a need to:

- Automate maintenance scheduling within ERP for a more refined production schedule;
- Upload up-to-date resource allocation information from the control systems;
- Upload equipment usage from the control systems to proactively schedule maintenance;
- Manage lab samples and map their results to production batches.

Interest in the integration of process control systems with enterprise systems results from the long-term goal to have only one single source of information

for entities like bill of materials, inventory, and genealogy. In addition, further advantages would be:

- Minimization of manual data entry errors;
- Minimization of recipe and Bill-Of-Material maintenance;
- Production of complex data summaries not available manually;
- Automated handling of process control system batch scheduling queues;
- Corporate benefits such as accurate planning, reduced inventories, accurate production cost tracking, accurate material tracking, optimal utilization of resources and people, accurate record keeping for quality certification, as well as specification of production targets.

4.2

Ethernet-Based Device-Level Networks

Ethernet TCP/IP based network technology is widely used in office environments to connect everything from mainframe computers to desktop PCs and printers. Although the Ethernet standard used on plant floors is the same as used in office environments, its implementation is considerably different, especially when it comes to redundancy. Several control system manufacturers are using Ethernet to connect controllers and operator interface devices together to form control-centric Ethernet architectures. Pushing nearer to the plant floor, a few manufacturers are forming device-centric Ethernet architectures where I/O devices are directly connected to an Ethernet network or are collected in Ethernet-compatible I/O data concentrators. The use of Ethernet in future projects depends on the application requirements, including how devices receive power and if intrinsic safety is needed. For either of those requirements Ethernet is not well suited. In addition, any factory level control network must be segmented and protected from business traffic to ensure the required reliability and determinism.

4.3

A Trend in the Future?

As pointed out in this review there is a wide gap between industrial requirements and academic research activities. On the academic side, sophisticated self-adapting controllers and complex mechanistic models to observe a process, on the industrial side, simple PID-algorithms, plenty of formalisms and process control based on empirical cultivation techniques. The time should come to merge the two approaches, in order to fulfill all the requirements of future biotechnological processes. The necessary computing performance is now available.

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