

The importance and future of biochemical engineering

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Abstract

Today's Biochemical Engineer may contribute to advances in a wide range of technical areas. The recent Biochemical and Molecular Engineering XXI conference focused on "The Next Generation of Biochemical and Molecular Engineering: The role of emerging technologies in tomorrow's products and processes". On the basis of topical discussions at this conference, this perspective synthesizes one vision on where investment in research areas is needed for biotechnology to continue contributing to some of the world's grand challenges.

KEYWORDS

Biochemical synthesis, bioprocess development, biomolecular engineering, individualized medicine, non-traditional organisms, synthetic biology

assembly; and the use of alternative biomanufacturing paradigms for enhancing biological conversion processes, such as microbial consortia, designed co-cultures, or cell-free systems. Other thematic areas include: bioprocess development for individualized medicine, forward-engineering for cellular control and predictable cell behaviors, which includes data-driven machine learning approaches for accelerating design, and engineering to understand & exploit new biology.

The topical areas listed below are by no means a comprehensive portrait of all current activities by biochemical engineers, nor is this the only current technical roadmap (e.g., <https://roadmap.ebrc.org/>). Rather, this Perspective is meant to synthesize one possible vision on where investment in research areas is needed for biotechnology to continue contributing to some of the world's grand challenges.

2 | THEMATIC AREAS

2.1 | Novel products and nontraditional organisms

Much of our view of biology and what is possible in biotechnology is shaped by what we learn in a small collection of well-characterized model cells like *E. coli*, *S. cerevisiae*, and CHO cells. Most educational resources are based on the discoveries made in these systems, and thus our view of life is often viewed in the context of these cells. Therefore, the fields of metabolic engineering and synthetic biology frequently turn to this short list of model cells as “chassis” for technology development. This has led to fantastic accomplishments, with undoubtedly great new advances in the horizon.

By contrast, investigation of non-model organisms, development of genetic tools in non-model organisms, and development of non-model organisms for use as chassis has been more limited. There are

many important reasons why we need to expand applied research activities with non-model cells and organisms (Figure 1).

- Alternative cells provide new opportunities for metabolic engineering and synthetic biology. Non-model cells may serve as superior “chassis” organisms as they can thrive in extreme environments and are already evolved for optimized performance of various capabilities. Non-model cells can provide different capabilities like stress-tolerant phenotypes and enhanced catabolic breadth (described in a recent review; Thorwall, Schwartz, Chartron, and Wheeldon (2020). Thus, alternative chassis may prove to be more suitable for future applications, including the use of cell-free systems (Silverman, Karim, & Jewett, 2019).
- Non-model organisms are already involved in a wide variety of well-established and scaled bioprocesses like wastewater treatment, metal mining, nitrogen fixation, and food production. Further investigation into the organisms found in existing bioprocesses will lead to new understandings of critical mechanisms, metabolic capacities, microbial competition, and mechanisms for robustness of cell–cell communication networks.
- Advances in biotechnology often arise from advances in basic biology, and important insights have been gained from non-model cells. Classic examples include restriction endonucleases and polymerases from thermotolerant extremophiles (Frock & Kelly, 2012). A more recent example is the discovery and engineering of a poly-ethylene terephthalate (PET) plastic degrading pathway found in a bacterium isolated from a bottle recycling facility (Yoshida et al., 2016). It is likely that new genomes and metagenomic sequence information from unculturable microbes and viruses in extreme and unusual environments can enable discovery of new biological capabilities and inspire new biochemical technologies.

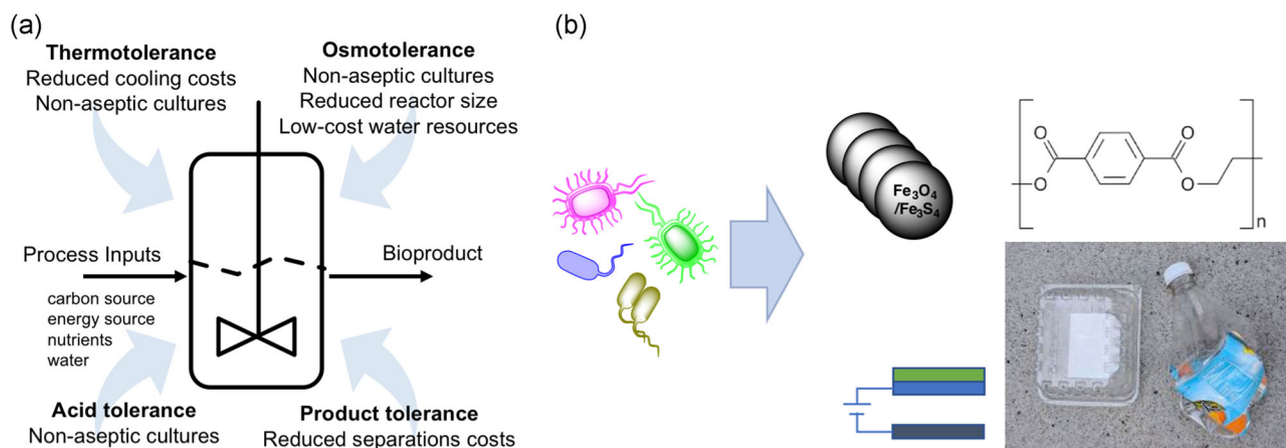


FIGURE 1 Novel traits in nonconventional microbial hosts can be exploited to create a new generation of biochemical processes. (a) Many nonconventional fungi and bacteria exhibit high tolerance to various environmental stresses that can occur during bioprocessing. Matching stress tolerant traits with critical bioprocessing challenges can save process costs and enable new designs that enhance product titer, rate, and yield. (b) Nonconventional hosts can be exploited for nonconventional processes like formation of magnet nanoparticles, bioelectrosynthesis, and valorization of plastic waste streams [Color figure can be viewed at wileyonlinelibrary.com]

A critical future goal of the biochemical and molecular engineering community will be the investigation, development, and engineering of non-model cells, components, and processes. This requires advances in computational tools for pathway prediction and large-scale systems biology data analysis to enable forward engineering. Such advances and research focus would especially benefit biotechnologies on the horizon such as biological/computer interfaces, waste recycling, and extra-terrestrial exploration (see a partial list in Table 2). As synthetic biology further expands into new organisms and microbial ecosystems it will be critical to replicate and even expand the biosafety strategies that have been used in the development of the classic model cells. There has already been interest in introducing biocontainment features into future generations of engineered cells (J. W. Lee, Chan, Slomovic, & Collins, 2018). In the sections that follow, we consider the near-future biotechnologies of sustainable protein production, and biological valorization of waste streams.

2.1.1 | Valorization of waste streams

Streams from municipal, agricultural, food, and plastic waste materials constitute a burden for communities, industries, nations, climate change and the environment more broadly. Increasingly, such streams are also viewed as an opportunity for utilizing the enormous quantities of chemical energy stored within them (Tuck, Pérez, Horváth, Sheldon, & Poliakoff, 2012). Many of these streams will be eventually converted to the greenhouse gases methane and CO₂ (e.g., in solid-waste disposal facilities or anaerobic wastewater treatment facilities) with very low, or zero, capture efficiency.

Generation of methane (biogas) from waste streams involves semisolid or liquid-stream methanogenic anaerobic digestion, largely based on the development of natural microbial consortia. Such processes are slow, not very effective, and thus not widely adopted. Challenges of producing fuels and chemicals from diverse feedstocks include the necessity of expensive biomass hydrolysis for effective fermentation, the loss of significant electrons generated from substrate catabolism to H₂, and extensive CO₂ loss due to decarboxylation of pyruvate to produce acetyl-CoA, the key starting intermediate for the production of most chemicals and fuels.

The ability to simultaneously use biomass substrates and gaseous substrates (renewable H₂ or syngas from various sources, such as from gasification of municipal or agricultural wastes) is of major technological significance as it would result in exceptional levels of substrate-carbon and electron utilization thus leading to high product yields. There are opportunities for combining biological and non-biological (e.g., catalytic/electrocatalytic) processes to achieve this goal. Technologies for utilizing both solid/semisolid and gaseous waste streams are therefore of major interest and should be the target of additional research investment.

In certain respects, valorization of plastic waste is an easier problem because waste is concentrated through commercial recycling operations with reasonable batch consistency. Biological conversion and upgrading of polyester and polyurethane waste plastic streams is particularly attractive because (a) ester and urethane bonds are accessible by enzymes; (b) plastic waste is much cheaper on a per mass basis than most existing carbohydrate feedstocks; (c) biological conversion routes are compatible with typical contaminants in plastic waste streams; and (d) monomers have similar reducing equivalents with current feedstocks. For example, the PET monomer ethylene terephthalate (C₄H₈O₄) has the same degree

Desirable phenotype	
Bacteria	
<i>Halomonas campaniensis</i>	Thermo-, osmo-, and alkaline tolerance
<i>Clostridium thermocellum</i>	Thermotolerance; lignocellulosic biomass breakdown
<i>Clostridium spec.</i>	Use of CO/CO ₂ as sole carbon sources
<i>Methanotrophs</i>	Use of gaseous alkanes as sole carbon sources
<i>Pseudomonas putida</i>	Solvent tolerant; catabolism of aromatics
<i>Acidothiobacillus ferrooxidans</i>	Acid tolerant; extracellular electron transfer
<i>Shewanella oneidensis</i>	Extracellular electron transfer
Yeast and fungi	
<i>Kluyveromyces marxianus</i>	Acid and thermotolerance; rapid growth
<i>Issatchenkia orientalis</i>	Acid and thermotolerance
<i>Yarrowia lipolytica</i>	Lipid catabolism
<i>Pichia pastoris</i>	Heterologous protein expression
<i>Neocallimastigomycota</i>	Lignocellulosic biomass breakdown
Cell-free systems	
Platforms	High-yielding, cost-effective, scalable bacterial systems for probing cellular function and biomanufacturing (<i>Escherichia coli</i> , <i>Vibrio natriegens</i> , <i>Streptomyces sp.</i> , clostridia, CHO, yeast, <i>P. pastoris</i> , plants)

TABLE 2 Selected nonconventional microbial hosts and cell-free systems for next generation bioprocessing

of reduction as glucose. Specific biochemical engineering challenges include developing enzymes that can efficiently deconstruct plastics to constituent monomers, and designing non-model organisms that can catabolize plastic monomers while also withstanding the necessary processing conditions. Additional challenges include a distributed “supply chain” and heterogeneity of contaminants in the waste streams. This will be a fertile ground for bioprocess engineers, protein engineers, synthetic biologists, and metabolic engineers.

The pressing environmental implications, and the need to move forward the concept of circular economy make it imperative that new thinking, new players and new investments are necessary to enable high-end and efficient processes to solve a problem of enormous global importance.

2.1.2 | Biochemical engineering opportunities in food and beverage production

Biochemical engineering has a long and storied history of supplying innovation for the food and beverage industry, including large-scale cultivation of microorganisms for nutrition. This development of such “single cell protein” was winding down as a research area before several authors on this perspective were born (Solomons & Litchfield, 1983). However, a resurgence of this topical area is led by commercialization of plant-based and cell-based meat products palatable to the end consumer.

As an example, the most publicized ingredient in the Impossible burger is genetically modified *Pichia pastoris* protein-rich extract containing a legume heme protein; when formulated in the burger this ingredient adds reddish color and flavor. This unapologetic use of genetically modified microorganisms opens the door for biochemical innovation in food products. Engineering microbial proteins that are more nutritious and yet still mimic the mouth feel of meat, or that can taste like sugar (Ming & Hellekant, 1994), or designing microbes with distinct flavor profiles tailored by metabolic engineering (Denby et al., 2018) are examples of innovations needed on the cellular engineering side. While large-scale fermentation processes for food and beverages exist, scale-up and bioprocess challenges for microbe-based protein are daunting: supplanting even 1% of U.S. daily protein consumption by single cell protein would require 750 metric tons of cells per day. More efficient cell harvesting and dewatering unit operations, programmed cell lysis, bioreactor design, and use of alternative feedstocks will be necessary before widespread deployment occurs.

The same rationale is valid for application as single-cell protein in present-day aquaculture. While aquaculture is the most-efficient and fastest growing protein generator for human consumption, one of its most relevant feedstocks is fishmeal which is limited in supply due to overfishing and therefore significantly compromises future sustainability of the aquaculture industry. Single-cell protein tailored to the specific needs of farmed fish and crustacean species might offer a solution.

Cultivated meat, by contrast, involves the in vitro production of cells present in meat used for human consumption. The cells used to produce cultivated meat include cell types present in meat such as skeletal myocytes and adipocytes from the mammalian, avian, and piscine cell lines of any meat-harvested species (E. A. Specht, Welch, Clayton, & Lagally, 2018). Recently, the National Academies of Science, Engineering, and Medicine (2017) noted the high growth potential of cultivated meat and identified it as an emerging biotechnology area. Efforts to achieve commercialization within the decade will require considerable attention to scale-up and large-scale manufacturing (M. J. Post, 2012). These practices include cell line selection and development, scaffolding, bioreactor design, cell culture medium optimization, and management of supply chain and distribution. One of the dominant barriers for cultivated meat to reach competitive prices with conventional meat is the cost of cell-culture media (National Academies of Sciences & Medicine, 2017). Traditionally, cell-culture media incorporated serum to promote cell growth, via the action of growth factors and other often non-defined components. Although serum-free and animal-origin-free media are able to support cell survival, proliferation, and differentiation (M. Post & van der Weele, 2014), a drastic cost reduction of both the basal medium and the growth factors would be required for economic viability at scale (L. Specht, 2020). Efforts directed towards drastically reducing the amount of growth factors needed, or the production of these factors in recombinant organisms, or the development of cheap protein mimotopes of these growth factors could offer a way out of this challenge. Metabolic modeling also offers an attractive avenue for benchmarking different ways of formulating a growth medium using either defined ingredients-only or supplemented with cell extracts (i.e., yeast or microalgae; Sathasivam, Radhakrishnan, Hashem, & Abd_Allah, 2019).

2.2 | Pushing past the limits of biochemical synthesis

2.2.1 | Combining chemical catalysis with biochemical conversion

Whenever the production of a new complex molecule is required from a given precursor there exists significant creative tension between chemists and biochemical engineers. Chemistry offers advantages in throughput, toxic intermediate tolerance, freedom to operate at high temperatures and the ability to leverage an existing chemical processing infrastructure. In contrast, biology allows for simpler processes, self-regulated pathways, making chemical changes in specific locations even for highly functionalized molecules. The recent review article by G.-M. Lin, Warden-Rothman, and Voigt (2019) highlights many of the new advances and remaining challenges. It is worth noting that continuous progress over the last few decades toward expanding the utility of enzymes, including advances in protein engineering, artificial enzyme development, and

high-throughput screening have opened new opportunities for chemoenzymatic synthesis in both aqueous and nonaqueous media.

While there are famous examples where both chemical catalysis and biochemistry were brought to bear (Anbarasan et al., 2012; Karp et al., 2017; Paddon et al., 2013), generally the two modes of production are deployed in isolation of one another. A number of retrosynthetic algorithms are available (Campodonico, Andrews, Asenjo, Palsson, & Feist, 2014; Henry, Broadbelt, & Hatzimanikatis, 2010; Kumar, Wang, Ng, & Maranas, 2018) for identifying a sequence of steps to a product using both existing and novel enzymatic steps. At the same time rapid progress has been made for chemical synthesis using rules-based pathway design (Klucznik et al., 2018). What is lacking is an integrated workflow for making decisions as to what steps will be carried out through biochemical conversions and which steps will be left to chemical catalysis (Wheeldon, Christopher, & Blanch, 2017).

How can we harness both chemistry and biology to produce previously unobtainable molecules? One potential new direction is the use of cell-free systems to create hybrid molecule products composed of elements derived from both chemical and biological synthesis strategies in the absence of viability constraints (Swartz, 2012). In another direction, repurposing the translation apparatus (including the ribosome and the associated factors needed for polymerization) to make sequence defined polymers comprised of novel monomers could lead to new classes of materials of defined atomic sequence, exact monodisperse length, and programmed stereochemistry. For example, synthesis of polyamides (outside of polypeptides) or aramid polymers could open new opportunities at the intersection of materials science and synthetic biology (Ad et al., 2019; J. Lee et al., 2019).

2.2.2 | Dynamic spatial assembly of metabolons and metabolic pathways

The design and assembly of so-called metabolons (structural-metabolic cellular complexes) and organelles mimics one of nature's strategies for maximizing productivity and carbon flux through biochemical pathways, and is a rich area of research for biochemical and biomolecular engineers. Metabolons or metabolosomes are multi-enzyme complexes that allow the direct passage of a product from one enzymatic reaction to a consecutive enzyme in a metabolic pathway, which in some cases may benefit from substrate channeling (e.g., when a side reaction competes for an intermediate in the bulk or an inhibitor is present that interferes with a reaction step; Wheeldon et al., 2016). Coordinated assembly and disassembly of these metabolons is an important factor in optimizing production of the desired metabolites. Natural organelle engineering has been effective in clustering key groups of enzymes—in peroxisomes and carboxysomes—and biochemical pathways believed to capitalize to at least some degree on enzyme localization and/or sequestration include tryptophan synthesis, the citric acid cycle, glycolysis, and purine synthesis.

The engineering concepts and physicochemical processes underlying the function of metabolons represent a scaled-down version of classical reaction engineering, and biochemical engineers have already made important contributions in modeling the behavior of systems ranging from one-dimensional scaffolds to three-dimensional microcompartments on multiple scales. Substrate channeling (Wheeldon et al., 2016), enzyme clustering (Castellana et al., 2014), and bacterial microcompartments (Jakobson, Tullman-Ercek, Slininger, & Mangan, 2017) have been the subjects of excellent modeling work, and these studies have provided important mechanistic insights and identified design criteria under which biochemical pathways will benefit from proximity and encapsulation effects. However, there are relatively few direct comparisons between such models and experimental systems, in part because well-characterized, precisely controlled experimental systems remain difficult to come by. Developing better techniques and methods to effect and control the assembly of scaffolded and compartmentalized systems both *in vitro* and *in vivo* is an exciting opportunity at the frontier of biomolecular engineering and related fields.

Many questions and challenges surrounding synthetic metabolons and organelles remain to be addressed, and several that emerge from the literature (Castellana et al., 2014; Jakobson et al., 2017; Kerfeld, Aussignargues, Zarzycki, Cai, & Sutter, 2018; Wheeldon et al., 2016) include the following:

- Controlling transport of substrates and products across the compartment shell/membrane.
- Predicting the membrane permeability of a given small molecule metabolite.
- Precisely controlling the number and location of encapsulated proteins.
- Harnessing experimental methods to analyze the physical configuration and molecular organization of the metabolon.
- Quantifying the kinetic effects of enzyme clustering and compartmentalization.

The new fundamental knowledge of how nature optimizes the productivity of biochemical pathways, together with the opportunities that such knowledge will afford for optimally engineering new pathways of practical interest, combine to make this area very fertile terrain for biochemical engineers.

2.2.3 | Microbial consortia and co-cultures

Many challenges in industrial biotechnology can be tackled by organizing microorganisms as “directed” consortia or even more well-defined “microconsortia”, such as synthetic co-cultures. These systems can be engineered using a more traditional top-down approach wherein microbe rich feedstocks are interrogated, prodded, and selected for specific purposes (Figure 2; Gilmore et al., 2019). Genomics-based methodologies and modeling are now being developed for the functional identification of the most useful consortia

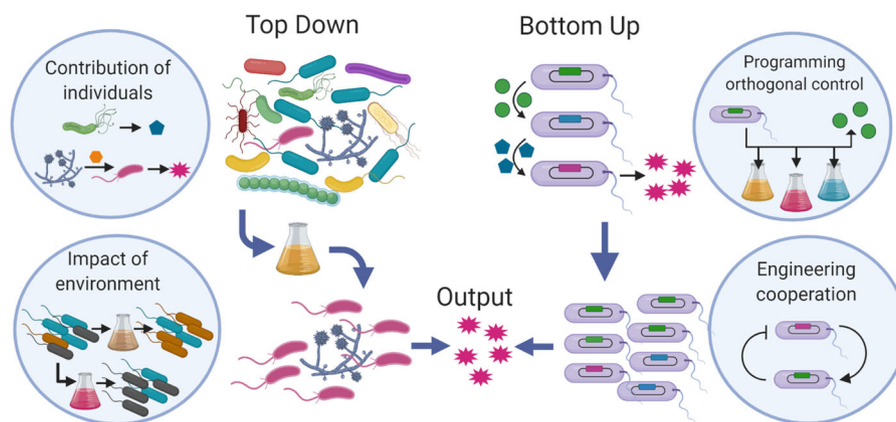


FIGURE 2 Microbial consortia or “microconsortia” can be designed using top-down or bottom-up approaches. In top down approaches, consortia exhibiting desired properties are obtained from natural environments and tuned or directed for the desired function or output. This approach would benefit from a better understanding of the contribution of individuals in the original consortia and environment, as well as a better understanding of how the environment affects the consortia composition and function. An alternate approach to designing mini-consortia uses bottom-up strategies. Here, individual strains or species are engineered to perform specific functions that are part of a larger task. In this approach, tools or strategies to guarantee the behavior of the individual strains despite changing or unknown environmental conditions are needed. Further, methods to engineer communication and feedback between strains could allow for maintenance of the consortia composition and function over time. This figure was created with [BioRender.com](https://www.biorender.com) [Color figure can be viewed at [wileyonlinelibrary.com](https://onlinelibrary.wiley.com)]

(Zuñiga et al., 2019), where the molecular bases for their intended functions are revealed and maintained. Importantly, complex initial sources, such as from anaerobic environments, can be accommodated (Solomon et al., 2016). Then, by using methodologies that reveal useful components for synthesis (Haitjema et al., 2017), “tuned” consortia might then be placed into processing environments for production. In this way, biomass feedstocks, particularly those that might otherwise be agricultural or municipal wastes, can be turned into useful, high value products. A major challenge to address for these applications is to maintain the consortia, or specifically, the precise composition of microbes (e.g., bacteria, fungi, and protozoans) that is needed to carry out the specific function, particularly if the processing conditions require extended time periods in industrial (nonnative) environments where population instability is well-known.

In these situations, a bottom-up approach may be more advantageous (Figure 2). In this scenario, co-cultures or other “mini consortia” can be assembled of sets of engineered cells forming highly functional cell systems that are programmed to execute specific tasks (Bittihn, Din, Tsimring, & Hasty, 2018; Jones et al., 2016; Lindemann et al., 2016; Shong, Diaz, & Collins, 2012). Additional design space is available for such systems relative to a monoculture engineered to perform the same task; each cell or strain can be optimally designed for executing a particular part of an overall task. In turn, the distribution of engineered cell subpopulations provides additional flexibility in the overall process design. For example, a hypothetical production process may be distributed among three cell types: one that employs raw materials and makes an intermediate, a second strain may also use a raw material, but also uses the intermediate synthesized by the first population to make a second intermediate, and the third strain might finish the overall process. The relative numbers of the three

strains can then be a control variable that is manipulated to ensure efficient production overall.

In both top-down and bottom-up situations, methodologies to coordinate subpopulation dynamics will be needed. These might involve external process inputs such as the addition of an inducer, an adjustment in oxygen or pH, or perhaps even process vessels that allow for fluid segregation or differential mixing. Conversely, in another novel approach, subpopulation dynamics could be created by rewiring native molecular communication systems like quorum sensing to autonomously control composition (Stephens, Pozo, Tsao, Hauk, & Bentley, 2019).

Specifically, new methodologies that recognize and interrogate the interplay between the external microenvironment and cell physiology will yield new insight on how to control cell behavior, particularly cell behavior that changes due to context. A cell's response, for example, to a molecular cue might be completely different depending on the redox potential in its microenvironment or on the identity of the neighboring cells. For example, in the human microbiome environmental factors, for example, chemicals, diets, etc. are known to impact the genotype-phenotype relationship and the development of diseases (Go, Nguyen, Harris, & Paul Lee, 2005). Thus far, they have been studied mostly for their involvement in metabolism (Sadler et al., 2018; Srivastava & Chan, 2008), signaling (and regulatory mechanism; Yang & Chan, 2009), and even biophysical interactions (Cho et al., 2019). However, it is becoming increasingly apparent that diets and environmental factors alter the microbiome (Lewis et al., 2015) as well as the epigenetic landscape (Cowley et al., 2012; Herceg, 2007) via DNA methylation patterns or histone tails to modulate the activity of genes and drive the development of disease. To investigate these new mechanisms, novel computational tools are needed to (a) decipher the microbiome and microbial

communities (Kim, Koh, & Rho, 2015) and how they impact the environment (diet)-gene-phenotype and (b) integrate data from the genetic, epigenetic, transcriptional, posttranscriptional, and metabolic levels and their interaction with the microbiota in the development of diseases. The differences between anaerobic, microaerobic and aerobic physiologies are well known, but are these conditions purposely manipulated to guide behavior? How are signal molecules perceived at the molecular level and how can we design consortia or guide microbiomes to adapt to and utilize cues to assemble valuable behaviors, synthesize valuable compounds, or degrade xenobiotics or other problematic compounds, or even guide human health? With additional tools that enable predictive biology and that exploit external inputs, we might better control systems that are comprised of microbiomes or consortia in a variety of places, not just in human locales, but in the rhizosphere and fresh or saltwater environments. Efforts in these areas are ripe for the talents of biochemical engineers who want to build on their strengths to address challenging problems that are sure to have a great impact on human health and our society.

2.3 | Bioprocess development for individualized medicine

Individualized medicine heralded a breakthrough when the Food and Drug Administration (FDA) approved Kymriah (Dolgin, 2017), the first CAR T cell immunotherapy and the first gene therapy in the United States. Following closely on the heels of cell-based gene therapies, directly administered viral vector-based gene therapy Luxturna for the treatment of a monogenic inherited vision loss disorder was approved by the FDA in 2018 (Food & Administration, 2017). Currently in 2020, there are 17 FDA-approved cell and gene therapies (<https://www.fda.gov/vaccines-blood-biologics/cellular-gene-therapy-products/approved-cellular-and-gene-therapy-products>), with further growth in this sector expected in the next decades. Cell and viral vector cell production for personalized medicine constitutes new challenges and opportunities for bioprocess engineers. In conventional bioprocessing, biomolecules are typically produced in stirred tanks that can be scaled up to meet demand. In the case of personalized medicine, particularly for autologous cell products, the challenge becomes scaling out production because each patient requires their own bioreactor. In many ongoing clinical trials, cell production is also decentralized and labor intensive: clinical teams at hospitals handle in-hospital cell manufacturing, often using batch cultures with little monitoring of cell culture variables such as cell density, pH, partial pressure of oxygen, and nutrient consumption rates. These process variables, when monitored, are often done off-line using sporadic culture sampling. Manual handling of cell therapy products using functionally open cell culture systems such as T-flasks remains commonplace. More automated systems are available, such as those utilized for autologous adoptive immunotherapies (Harrison, Ruck, Medcalf, & Rafiq, 2017; Iyer, Bowles, Kim, & Dulgar-Tulloch, 2018), but even these have limited on-line monitoring and feedback control over

cell culture parameters. Automation and regulatory requirements to minimize risks of contamination as well as product variability create a strong drive towards the use of closed cell culture systems such as cell culture bags. As most preclinical studies are conducted in polystyrene vessels, the transition to bag-based cultures can lead to changes in cell-surface interactions and other culture parameters such as gas exchange (Fekete, Béland, Campbell, Clark, & Hoesli, 2018). There is a strong need to use scale-down culture systems during preclinical development which better reflect manufacturing methods and culture vessels at clinical scale.

For allogeneic cell products, scale-up can be performed and can rely on bioreactor designs that approach more conventional biomanufacturing. However, the challenge of on-line monitoring of a cell-based product remains. Moreover, many allogeneic cell therapy products such as mesenchymal stem cells or induced pluripotent stem cell-derived products are anchorage-dependent cells. Scale-up thus often relies on increasing the surface area for cell adhesion, for example, using microcarriers, hollow fiber bioreactors or stacked vessels—increasing the complexity of automated handling.

Viral vector production—whether for transduction of cells *ex vivo* or *in vivo*—at clinical scales with high reproducibility also remains challenging (McCarron, Donnelley, McIntyre, & Parsons, 2016). Many research-scale viral vector production system utilize anchorage-dependent cells which require hollow fiber bioreactor or microcarrier systems which are much more complex to scale up. With cell lines adapted to suspension culture such as human embryonic kidney cells, process intensification is an area of focus. Productivity does not only require high yields of viral particles, but also of properly assembled viral particles that maintain their functional capacity to transduce and express transgenes in target cells. In-line or rapid off-line monitoring of viral particles would significantly accelerate upstream process optimization. Finally, novel downstream purification methods that are scalable and that can resolve functional from nonfunctional viral particles are needed.

Although there have been significant advances in adapting culture systems to challenging cell therapy products over recent years, some of the practical questions that need to be addressed are:

- Can we formulate a list of overarching cell culture parameter ranges required for cell and therapy products in adherent versus suspension culture?
- What biomaterial approaches or genetic engineering methods may we employ to control the homogeneity of the desired cell populations?
- How can we make current culture systems more flexible and adaptable by end-users (including clinical centers) to facilitate manufacturing of several cell therapy products with a single system?
- What in-line methods could we employ to better assess and control cell and gene product quality?

Cellular therapy is set to revolutionize the treatment of cancer and conditions where small molecules and other biologics have not led

to a cure to date. The growing list of approved cell therapy products (<https://www.fda.gov/vaccines-blood-biologics/cellular-gene-therapy-products/approved-cellular-and-gene-therapy-products>) not only for people suffering from blood disorders and cancers, but also for cartilage, retinal and other tissue defects portends a new era in the treatment of degenerative disease. Groundbreaking clinical trials are testing the safety and efficacy of embryonic stem cell-derived products transplanted in various encapsulation devices to treat type 1 diabetes (Moeun et al., 2019). Addressing the bioprocessing challenges listed above is critical in assuring the safety, efficacy and accessibility of these life-saving products.

2.4 | Forward engineering for cellular and biomolecular control

2.4.1 | Integration of mechanistic based models with data driven approaches for protein- and cell-based engineering

Since the advent of the biochemical engineering discipline mechanistic models based on kinetics and thermodynamic constraints have guided experiments. We now have a torrent of high quality data from myriad omics technologies, deep mutational experiments of protein and RNA-encoding sequences (Kowalsky et al., 2015), and facile high-throughput strain development in many organisms spurred in part by the CRISPR revolution (Schwartz, Hussain, Blenner, & Wheelton, 2016). To what extent could these new large data sets, with potential for more modern machine learning approaches, enhance current modeling techniques? Compared with current models, what kind of biological knowledge could we gain by using machine learning?

An illustrative example comes from protein science. The protein folding problem is typically formulated as predicting an accurate atomic structure of a protein given its sequence of amino acids. In 2018, the winners of the blind prediction CASP challenge were a group of Alphabet engineers without specific training in this area. The team, dubbed AlphaFold, outperformed all other scientific groups in the world and really advanced the field by about 2–3 years (AlQuraishi, 2019). Importantly, they used the mechanistic insight that positions that are close in distance tend to co-evolve together. This insight is not new and has been developed in the literature over the past two decades (Morcos et al., 2011). They were successful in large part because the existing data sets of tens of millions of accurate protein sequences and over a hundred thousand protein structures were vast, centralized, and curated. They used deep learning to learn a differentiable potential between co-evolving residues that is specific for each protein.

This example is particularly instructive because it tells us a few things about how our community should approach this opportunity. First, we want good data and heaps of it, no matter the source. Methodological advances should be encouraged for collecting large amounts of phenotypic and genotypic data on engineered strains and

activities and biophysical properties of proteins. Similarly, strong efforts to centralize already existing literature data sets should be supported, perhaps as a community effort. As an example, the protein engineering field now does this with ProtaBank (Wang et al., 2018). Second, the AlphaFold team improved on existing mechanistic insights into how coevolution of residues predicts distance in the folded polypeptide chain using their deep learning approach. They also used an ensemble model with existing structure-based prediction using physically realistic potentials in the macromolecular modeling software package Rosetta. The field should embrace ensemble models and related techniques may be applied to nail down the thermodynamic driving forces for resolving kinetics of intracellular fluxes (Gopalakrishnan, Dash, & Maranas, 2019) or better use of evolutionary and/or coevolutionary networks and other mechanistic insights to engineer stability in enzymes (Ritter & Hackel, 2019). Here is where deep learning may be particularly useful in identifying very strong mechanistic bases for why outcomes look the way they do, given a range of potential inputs. Third, the AlphaFold team originally looked at much more complicated machine learning models using features that do not have such mechanistic insight, which they discarded because of the strength of the simpler and more powerful coevolutionary analysis. Simpler features grounded in physicochemical or evolutionary mechanisms will ultimately be more useful, more likely to lead to biological insights that can be exploited, as most of what we do is grounded with strong constraints set by physical chemistry.

Finally, we should be realistic about the data we have and can generate. Existing linear and nonlinear regression based models work well in a variety of contexts. For example, one of us (T.A.W., unpublished) has found in protein engineering that linear regression seems to work fairly well for prediction of protein activity, consistent with reports from more limited data sets (Fox et al., 2007). These simpler regression models also have the advantage of being more interpretable.

For cell engineering specifically, there are clear recommendations for efficiently exploring the vast genetic space to achieve actionable and or valuable cell engineering outcomes:

1. Mine existing data sets: Many large, unbiased genetic characterization studies have been conducted to date on model organisms and have been published. We need to leverage what has already been done to find patterns. This requires us to aggregate and organize the data sets across multiple studies and leverage searchable databases. It also requires a higher level of engagements/knowledge sharing from industry. Here community efforts to centralize such data sets, as mentioned above, should be strongly supported and encouraged. As an example, some studies have comprehensively tested the genetic landscape for host organisms (e.g., genetic transcription engineering). Can we retroactively review these studies and outcomes to learn what worked and perhaps why? Can we leverage those findings to understand how to effectively truncate a genetic search space without losing quality/positive outcomes?

2. In silico tools: Meticulous experimental studies are time and resource consuming. Search space is more efficiently managed using good in silico models. We need to continue to enhance metabolic models, and pressure test the quality of models that are developed using diverse metabolic pathways (i.e., not just central carbon metabolism). Going forward, there should be more emphasis on comprehensive, complex metabolic functions (glycosylation, lipids, polyphenols, etc.), which complements the complex products the field is now interested in producing using cellular hosts.
3. Understand what is host/cell line specific versus biologically universal: A lot of excellent studies are published on one cell line/host to understand or fix specific biology. We need to understand when these findings can be leveraged for a different cell line/product, and when we can avoid repeating cellular optimization/engineering efforts. To build this understanding, we should consider vertical organism testing, that is, progressing an optimization with a specific outcome in mind first through a single celled, prokaryotic organism, then through a single celled eukaryotic organism, and finally through a multi-celled eukaryotic cellular host.
4. Beware of model protein products! Proof of concept work on simple proteins may not translate to complex targets. It could mask/mislead/not scale to the desired, applications and products. We should incorporate this consideration into study designs to ensure the best quality information is captured.

2.4.2 | Transforming cellular control and predictable cell behaviors through synthetic biology

A major issue in biomanufacturing and bioprocessing is heterogeneity and lack of control in cell behavior manifesting in alterations of process parameters and product quality. We need to understand and control the sources and mechanisms of heterogeneity to achieve better process control, reproducibility and reliability. One way to address this challenge is to build orthogonal, tunable tools that operate on time scales faster than the process being controlled to make cells more readily manipulated and directed towards the generation of desirable products.

Engineering cellular systems with predictable behavior requires diversification of tools to achieve control at the molecular level. Current tools to control cell behavior are mainly based on transcriptional regulators and have been successfully evolved through a variety of protein engineering methods. There is a pressing need currently to identify new tools and new methods for identifying appropriate dynamic control elements to use in larger systems. Protein-mediated regulation typically operates over faster time scales than transcriptional and translational control and may be coupled directly to endogenous pathways and without the need for genomic integration (Budihardjo, Oliver, Lutter, Luo, & Wang, 1999), enabling dynamic control. Repurposed CRISPR-Cas molecules have also been explored (Xu

& Qi, 2019). Despite successful methods for exogenous control over CRISPR system, methods for internal controls remain a challenge. Efficient tools for tuning CRISPR activity, such as the recently discovered anti-CRISPRs, are needed for the future development of synthetic CRISPR-mediated circuits (Nakamura et al., 2019). Finally, naturally occurring epigenetic programs underlying cellular differentiation and development provide new opportunities for the design of control systems based on molecular writers and readers of chromatin signatures (Park, Patel, Keung, & Khalil, 2019).

Larger control systems can be assembled as more control elements are developed. Yet, there are many open operational questions for how cellular pathways detect and process input signals. First, the quantitative and dynamic input features that are perceived by natural and synthetic control systems are not always fully characterized for systems. It has become increasingly apparent that input dynamics rather than absolute values play significant roles in shaping the ultimate cellular outcome. Second, the system design needs to be carefully determined: extrapolating the design rules of classic microbial two-component systems to predict more complex signaling networks has proved to be a nontrivial endeavor, requiring tuning of control elements guided by deterministic and stochastic modeling carefully deployed to predict system behavior.

Predicting pathway behavior has proven to require quantitative modeling to develop an accurate understanding of even relatively simple systems (Ha & Ferrell, 2016). Ligand-controlled responses such as growth factor pathways, for instance, can respond to input concentrations with a diverse range of sensitivities, pointing to the critical need to build operational models of cellular systems based on quantitative descriptions of the input-output properties of each signaling pathway.

Critical recommendations related to progress in the development of cellular control systems include:

- Experiments should focus on single cell analyses to avoid confounding effects of population heterogeneity. Because cellular behaviors are often unsynchronized, it is also important to explore the dynamic response of single cells to avoid artifacts from static single cell or population measurements. Additionally, where possible, researchers should capitalize on gene-editing technology to reduce population heterogeneity.
- Studies of cellular control systems should rely on reconstitution of minimal versions of circuits and gene networks; isolation of minimal version of cellular pathways from natural inputs and outputs enables studying signal processing capabilities systematically and generating predictive models that recapitulate the governing features of different control networks.
- As larger scale genetic circuit engineering remains challenging, it is important to leverage the predictive power of mathematical modeling and integrate models and experiments to explore the behavior of complex cellular systems across parameter regimes.

2.5 | Engineering to understand and exploit new biology

2.5.1 | Building and exploiting interface between electronics and biology

Semiconductor technologies have transformed our abilities to access, store, process, and communicate information by enabling increasingly smaller, cheaper, more powerful, interconnected, and easier-to-use electronic devices. Synthetic biology will enable the extension of these modalities to interface with electronics—by rewiring and programming cellular processes that manipulate chemical information at the molecular scale, using redox as a vector of information transfer (Liu et al., 2017)—in ways that facilitate information exchange with electrodes (Tschirhart et al., 2017; VanArsdale et al., 2019). There has already been remarkable progress in spanning biological and electronic communications for an important subset of problems involving the ionic electrical modality, including advances in neuroprosthetics and in understanding and mapping brain function. *Molecularly* based information transfer and notably, redox enabled communication is widespread in biology: it is used by the immune system for inflammation and wound healing; it underpins communication within the gut, and potentially between the gut and brain; and it enables communication in the biosphere (e.g., cells in the plant roots can detect/respond to activities in its rhizosphere), to name a few. To enable redox-based communication, future opportunities lie in the fabrication of “smart” materials interfaces that integrate biological recognition and computation while facilitating information transfer to and from the devices at length and time scales that are often viewed as discordant.

There is tremendous potential for the development of devices that seamlessly transfer information to and within biology. To provide just one example, wearable devices such as smart watches that provide actual chemical information in addition to what is currently available (i.e., moisture, temperature, and cardiovascular function) will radically transform our everyday lives. New efforts in electron transfer, redox biology, materials and surface characterization and assembly, will be needed in addition to traditional expertise in mass and momentum transfer, reaction kinetics, and thermodynamics, to create effective systems for information transfer into and out of the biological system.

2.5.2 | The biology and biotechnology of extracellular vesicles (EVs)

EVs are membrane vesicles that carry RNAs, proteins, lipids, and sometimes DNA from their parent cells (Kao & Papoutsakis, 2019). EV generation takes place under cellular activation or stress. Cells use EVs to communicate with other cells by delivering signals through their content and surface proteins. Besides mammalian cells, outer membrane vesicles (OMVs; Anand & Chaudhuri, 2016), derived from Gram-negative bacteria, are involved in stress response,

promoting survival, pathogenesis, and interaction between bacteria in a community. Gram-positive bacteria generate a large number of EVs, as well, but their role in intercellular communication remains largely unexplored. Over the last few years, EVs have emerged as important mediators of intercellular communication regulating an ever-expanding range of biological processes, both on normo- and pathophysiology. The former includes enhancing and accelerating native developmental programs in immunology, vascular repair, and angiogenesis, while the latter include carcinogenesis and cancer metastasis, neurodegenerative disorders, and infectious and cardiovascular diseases.

On the basis of their currently known biology, EVs are suitable for a broad range of applications, from minimally invasive diagnostic applications to therapeutic interventions, including cell therapies and macromolecular drug delivery. In addition, there are two new emerging EV subfields. One is the role of microbial EVs in microbial consortia activities, including those of the microbiomes, and in the plant-to-microbe interactions. The other is based on the metabolic activities of EVs independently of the parent cells. The latter can be the basis for designing and employing efficient cell-free systems for advanced biocatalysis including combinatorial biosynthesis, but distinct from the current technologies that are based on *in vitro* transcription and translation.

Both EV cargo and membranes can be independently engineered and used for various applications (Kao & Papoutsakis, 2019). To pursue such applications involving EVs, better EV characterization, as well as better understanding of the mechanisms of cell targeting (Jiang, Kao, & Papoutsakis, 2017) and methods for EV biomanufacturing are needed. This is a relatively new field, especially regarding microbial EVs, but there is great potential in a broad spectrum of applications, thus making EV-funding investments a worthy cause.

2.6 | Perspective

Biochemical engineers are involved in solving many of the world's greatest challenges. This perspective synthesizes where research investment should be strengthened to enhance the impact by the discipline. For each thematic area there are clear recommendations moving forward.

First, further and more sustained investment and research is needed in developing efficient ways to build new genetic tools in non-model organisms. Novel products requiring non-model organisms or cell-free systems should be particularly supported. Additionally, novel technologies enabling microbial process scale-up and downstream processing are strongly desired.

Second, developing truly sustainable bioprocesses requires circumventions of current limitations on cellular biochemical synthesis. High on the list are methods or workflows to determine how to split a process between biochemical conversion and chemical catalysis. Cell-free systems creating hybrid chemical/biological synthesis is one approach to remove cellular constraints; continued development of

such systems should be supported. There are a number of fundamental questions on metabolons that can be addressed with careful experimentation. Finally, control mechanisms should be discovered and engineered for tailoring precise, stable, compositions of microbial consortia for various bioprocesses.

Third, several aspects of bioprocess development for individualized medicine need to be studied, including determining cell culture parameter ranges for adherent versus suspension cultures, improving the homogeneity of the cell populations, continued innovation for increasing flexibility and adaptability of cell culture systems, and developing better in-line methods for assessing and controlling product quality while assuring accessibility to these life-saving therapies.

A fourth thematic area in forward engineering for cellular engineering and biomolecular control already commands significant research support, which should continue, but with several clear recommendations. The current published deluge of high quality phenotypic and genotypic data on engineered strains and proteins should be centralized, perhaps as a community effort. Machine learning approaches to analyze, evaluate, and predict properties should be undergirded by evolutionary and/or physical chemistry principles. Researchers should be wary about using model cell lines and protein products to extract out biologically universal principles. Experiments with engineered networks should focus on single cell analyses as well as engineered homogeneous cell populations with robust mathematical modeling to guide understanding of the phenotypic parameter space.

The fifth and final thematic area involves engineering to understand and exploit new biology. Here new topical areas in merging electronics and biology and exploitation of EVs were discussed, along with the attendant challenges inherent in these new fields.

Progress on these thematic areas is necessary for solving grand challenges in environmental & energy sustainability, and the next generation of safe, effective medicines.

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CONFLICT OF INTERESTS

The authors declare that there are no conflict of interests.

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