Single-Use Systems As Principal Components in Bioproduction

William G. Whitford

Single-use systems (SUS) have become an accepted component of animal-cell–based bioproduction. No longer a merely exciting possibility, they have emerged as a significant and growing resource for companies to use from process development to manufacturing of approved products. Having been examined for years in less regulated environments, off-the-shelf SUS are now in regular use to some extent in nearly every segment of the production train by contract manufacturing organizations (CMOs) and biopharmaceutical companies in mid-scale production applications. For many operations, in fact, the question has evolved from, Is there an SUS available that can support my process? to, Which system best supports my needs?

Drivers of this rapid acceptance and widespread use in such a conservative industry have been well reviewed in recent years as early adopters experimented with emerging technologies and products (1). In general, advantages are found in capital investment, cost of goods, safety, production scheduling, surge capacity, and process replication. But this acceptance may not be more generally recognized. Hybrid or integrated implementation is common, with disposable elements operating in line or in concert with reusable materials. Also, bioproduction incorporates many divergent products and production processes, each with its own requirements. Although SUS are now fully accepted for some operations, their potential in others remains to be demonstrated.

Single-use technologies are in a state of remarkable growth, which determines a range of implementation maturity — from systems clearly established by industry leaders to emerging systems being introduced by equipment vendors and academic institutions. Even though such exciting innovations as entirely disposable operations and even closed and modular process trains are possible if yet to be fully realized, the ubiquitous existence and acceptance of an array of disposable components has been firmly established (2–4). Reviewed here are some previous concerns regarding SUS implementation that have been clarified recently (Table 1).

**Cost and Finance Advantages**

Although accurate comprehensive models of SUS implementation have turned out to be more complex than originally envisioned, solutions or clear paths forward are now available. To begin with, biopharmaceutical manufacturing costs and their modeling per se are not very perspicuous (5). Nevertheless, the consequences of SUS implementation to facility and operational costs — as well as financial strategies — have been determined for many applications and implementations (6). In fact, a growing number of models and case studies are publicly available and supplied for the most part by independent consultants.

Although the list of general features and advantages provided by disposables is long and growing (see “Demonstrated Advantages” box), the cost and financial implications of a particular system often remains case specific. Project- or facility-specific factors include such considerations as what conventional manufacturing equipment is currently in place; the desired type and timing of...
Significant study has led to the conclusion of equivalent or reduced current status. Continued development has increased flow rates, improved ease of use and reliability and added to the number individual technologies and product vendors. Multiple vendors support diverse products for many operations, and multiple technologies support distinct processes (e.g., mixing, sensing, connectivity).

Materials acceptability

SUS are now overcoming classification and scale-influenced qualification demands, including mechanical strength and leachables.

Process development and control

Supported by applicability of production-scale control in development, process formats comparable to conventional systems are available.

Technical transfer

Overlapping system capacity ranges with uniformity of component materials and configurations support transfer.

Table 1: Most initial challenges for SUS have been resolved by many available systems.

<table>
<thead>
<tr>
<th>Demand</th>
<th>Current Status</th>
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<tbody>
<tr>
<td>Cost and finance advantages</td>
<td>Rigorous modeling and a number of published studies delineate significant savings SUS provide for operations in many implementations.</td>
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<tr>
<td>Unit operations support</td>
<td>SUS exist for most steps in bioproduction. Many have seen industry acceptance or been used to manufacture products under FDA review and in clinical trials.</td>
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<tr>
<td>Products and platforms support</td>
<td>Success has been demonstrated in production of significant product categories and for all major production modes and formats.</td>
</tr>
<tr>
<td>Component offerings</td>
<td>Multiple vendors support diverse products for many operations, and multiple technologies support distinct processes (e.g., mixing, sensing, connectivity).</td>
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<td>Materials acceptability</td>
<td>SUS are now overcoming classification and scale-influenced qualification demands, including mechanical strength and leachables.</td>
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<td>Technical transfer</td>
<td>Overlapping system capacity ranges with uniformity of component materials and configurations support transfer.</td>
</tr>
<tr>
<td>Scale-up</td>
<td>Case studies have established practical comparability (e.g., mixing, flow, power/volume, and $k$,$\alpha$ explained through computational fluid dynamics as well as stochastic and empirical multiscale modeling).</td>
</tr>
<tr>
<td>Regulatory</td>
<td>FDA likes reduced qualification/validation demands and contamination potential, with increased design flexibility. SUS are used to make some approved products and many filings and trials.</td>
</tr>
<tr>
<td>Probes, sensors, and monitoring</td>
<td>SUS mixers, bioreactors, and columns support both classical and an increasing number of single-use sensors and samplers.</td>
</tr>
<tr>
<td>Connectivity</td>
<td>Continued development has increased flow rates, improved ease of use and reliability and added to the number individual technologies and product vendors.</td>
</tr>
<tr>
<td>Ecological footprint</td>
<td>Significant study has led to the conclusion of equivalent or reduced environmental impact over classical systems for many SUS in many implementations.</td>
</tr>
<tr>
<td>Performance</td>
<td>SUS show scalability, reliability, flexibility, and productivity reported to be equivalent with or superior to many traditional systems.</td>
</tr>
<tr>
<td>QbD and PAT</td>
<td>In general, SUS capabilities are equal to classical systems, but some unique advantages support increased process understanding.</td>
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For years, single-use technologies (with a few notable exceptions) have had a more restricted range of application in downstream operations. But many exciting recovery-through-finish systems are appearing based on single- and limited-use technology. One difference in the implementation of disposable technology in downstream (rather than upstream) processes is that purification requirements often determine more stringent specifications for material robustness and leachables. Also, surface binding characteristics and particulate burden must be considered. Due to the proximity of downstream disposable elements to final products, risk-analysis may indicate a heightened requirement for testing-based validation and documentation. SUS are now gaining acceptance for large-scale purification, including scalable tangential-flow and depth filtration units (Figure 1).

A surprising number of new, single-use, large-scale purification and polishing technologies are now appearing and at varying degrees of acceptance. The industry’s early adopters are regularly using prepacked, prequalified, and presanitized large-scale chromatography columns designed for limited- or single-use containing many popular resins (13). Scalable single- or limited-use membrane ion-exchange chromatography is now offered in both strong anion- and cation-exchange forms. Membrane adsorbers have been successfully applied to purify DNA, protein, viruses, and other high–molecular-weight products and for DNA and endotoxin removal in polishing steps.

One of the more powerful technologies now making an appearance involves simulated moving bed (SMB) chromatography. For example, the BioSMB system from Tarpon Biosystems (www.tarponbiosystems.com) comprises a series of continuous, disposable flow paths converting a batch chromatographic process into smaller multiple-separation-device processes. It even allows for maintenance of chromatographic resins and buffer systems already familiar to users (14).
Single-use technologies and products are also changing drug-product–based operations (15). Fill–finish operations requiring no laminar-flow controlled environments are now facilitated by closed-system and single-use bags, transfer manifolds, and connectors. Automated, aseptic filling of injectable drugs using SUS is supported by high-speed equipment that can accept ready-to-fill containers.

**Materials Acceptability**

Despite related concerns in biopharmaceutical production, disposable (plastic) materials in flasks, tubings, and filter membranes have been a part of biomanufacturing trains since the industry’s inception. But introduction of such large-scale systems as mixers, bioreactors, and separation/purification assemblies has required examination of new resins and castings, an expanded range of process conditions, and proximity to active pharmaceutical ingredients (APIs) and dosage forms. Even materials previously characterized and approved (e.g., USP Class VI or ISO 10933–1 compliant) for small-scale use need risk assessments of the greatly increased ratio of single-use surface to process fluid or product mass.

Individual system adopters, government agencies, and nongovernmental organizations have worked to establish clear and practical paths forward. For example, the Bio-Process Systems Alliance (BPSA, www.bpsalliance.org) has developed disposables “best practices” and technical guides on such topics as extractables and leachables testing (16). The International Society for Professional Engineering (ISPE) has established a Product Quality Lifecycle Implementation (PQLI) biotechnology task team to examine these materials issues from a QbD perspective (17). Topics currently in focus include establishing consensus among countries and standards-setting organizations: e.g., the US Food and Drug Administration (FDA), European Medicines Agency (EMA), International Standards Organisation (ISO), American Society of Mechanical Engineers (ASME), and pharmacopoeia.

**Scale-Up, Technical Transfer**

SUS that present consistent configuration and materials through a range of scales ease transfer from small-scale (development) to large-scale (manufacturing) operations (Figure 1). For some time, it has been possible to perform upstream process development in submilliliter-scale SUS microbioreactors. Recently it became possible to move from process optimization in a 3-L stirred-tank disposable bioreactor to production in a 2,000-L stirred-tank disposable reactor while maintaining the very same dissolved oxygen and pH probes, porting, and automated control system compliant with 21 CFR 11 (e.g., Delta V SIS). Published case studies report the remarkably robust scalability of SUS from micro- to production-scale reactors (18, 19).

**Demonstrated Advantages of SUS Over Classical Systems**

**Financial Advantages**

**Reduced Capital Investment**: Reduces service requirements, production process and classified area footprint; reduces facility design, install, and qualification activities; converts some up-front fixed costs to variable long-term; supports inexpensive establishment of efficient surge capacity

**Reduced COG**: Lowers utility costs; requires fewer manufacturing steps and labor requirements; reduces quality activities and preventative maintenance costs; lowers process flow and equipment modification costs; supports rapid and economical process replication and decentralization of production; provides cost distribution advantages

**Speed and Efficiency Advantages**

**Installation Ease**: Reduces plant development and start-up time and supports “hybrid” system installations and alterations

**Operational Ease**: Reduces process steps and validation steps and activity; eliminates cleaning operation costs and downtime; increases plant capacity and production scheduling flexibility, supports aggressive scheduling and campaigning; facilitates flexible approaches in process flow and layout; simplifies and accelerates product changeover and turnaround; supports an evolving “plug-and-play” approach to operational modules and enumerable hybrid reconfigurations; eases CGMP cold-chain logistics and good storage practices, with integral cryopreservation

**Change Ease**: Supports process modularity and flexibility; increases ease in reconfiguring and extending a production facility, with economical and rapid site-to-site operations transfer; provides for general equipment and process facility independence; supports economical system upgrades and advancements

**Environmental and Safety Advantages**

**Sustainability**: Reduces waste, effluent, and use of harmful cleaning materials; lowers water and power consumption, fouled water release; reduces carbon footprint

**Safety**: Reduces the number of activities open to contamination; offers a growing number of entirely closed operations; lessens operator exposure, lot and product cross-contamination risk; improves compliance values (reduces error potential); eases material transfer between diverse biosafety or classification levels; increases sterility assurance (irradiation rather than steam)

**Other Advantages**

**Manufacturing-Scale Performance**: Provides demonstrated scalability within available system sizes; supports existing process sensing, monitoring, modeling, and control

**Growing Technology**: Increasing numbers and capabilities of novel systems (e.g., connectivity, sensing, chromatography); promise of further improvements (processes and models in infancy, yet to reflect true potential); promise of an expanding range of “closed-system processing” across unit operations

**Growth in Acceptance/Validation**: Provides for expanding applications up- and downstream

**Figure 1**: Scalability of single-use components is demonstrated by disposable depth-filtration systems available in the same media from milliliter to 2,500-L capacities and beyond. Seen here is a Zeta Plus encapsulated system from 3M Purification Inc. (www.3mpurification.com).
Supported Products and Platforms
The utility of many disposable processing solutions has been demonstrated at production scales using most standard upstream and downstream platforms.

Upstream this includes fed-batch and some perfusion cell culture modes for all major production animal cell lines, from suspension Chinese hamster ovary (CHO) cells to adherent VERO cells to Sf9 insect cells with the baculovirus expression system (BVES). Processes have been established in single-use bioreactors for different growth rates and cell masses, including attached cultures using microcarriers. Success has been reported even with some microbes that have low growth rates or that generate low cell mass. The utility of single-use bioreactors has been demonstrated in production of proteins, viruses, nucleic acids, cell-based products, and vaccines (14).

Downstream operations are supported for many products. These involve the full range of applications from depth filtration and centrifugation to purification and polishing. For some particular processes, such as protein A chromatography of monoclonal antibodies (MAbs), the current single- or limited-use technologies and commercially available products can be uneconomical.

Regulatory Compliance
Steps toward meeting regulatory requirements for SUS have been encumbered by a paucity of standards, guidances, and best practices specific to bioprocessing disposables. For the most part, compliance has been established with an eye to standards established for traditional systems. However, not only have a number of processes involving SUS been validated in biopharmaceutical applications, but comprehensive validation guides and application-specific services are now provided by vendors to aid in efficient implementation of their products. Some are establishing the quality attributes of particular systems, providing the testing required for their qualification, and determining how the contents of regulatory filings might be orchestrated when SUS are involved. Furthermore, the FDA is reported to be very supportive in establishing such processes in the biopharmaceutical industry (20).

Probes, Sensors, and Sampling
First-generation monitoring approaches in SUS operations relied on the creative connectivity of classical probes to disposable equipment. Solutions involving existing aseptic quick-connects have both supported the new systems and provided a continuity in the transition to disposables. However, they have suffered from limited capability at smaller scales. Manufacturers are now working to establish a second generation of sensors, incorporating a number of novel and powerful technologies (21).

A growing number of commercially available probes even have integral and presterilized disposable product-contact surfaces. Included are probes for temperature, pH, and O\textsubscript{2}, CO\textsubscript{2}, pressure, and cell-mass. Many are compatible with Object Linking and Embedding (OLE) for Process Control (OPC) and compliant with 21 CFR part 11 digital automation systems. New sensing technologies being applied to SUS or SUS-compatible probes include fluorescence, capacitance, sensor-patch, and various infrared (IR) analyses (22). Cell-free sampling ports and interfaced autosamplers with demonstrated SUS utility support integration of the same instruments used in traditional glass and
Coupling and Transfer
A number of solutions are now commercially available to address the challenge of connectivity. From hybrid disposable-compatible connectors to fully disposable genderless connector systems to in-line welding, process designers have an increasing diverse pool of approaches from which to choose. Although the issue has been resolved for many implementations, challenges remain in regard to particular scales, flow rates, and process conditions.

Environmental Footprint
Initial reactions of many people are that systems reliant on “disposable plastic” components would be more detrimental to the environment than are classical systems. But on deeper examination, we find that many types of considerations arise.

First, SUS plastic waste must be compared with the many wastes generated by a classical system being replaced. Consider the fouled water disposal that follows clean-in-place (CIP) procedures and the fuel burned to generate steam for steam-in-place (SIP) systems.

Second, the distinct and divergent categories of “environmental stress” receive varying degrees of concern even from experts. For example, it is difficult to weigh the relative harm imposed by carbon generation against landfill use and water consumption (23).

A host of implementation-specific considerations include national, geographical, and economic imperatives introducing new factors such as regional standards on wastewater disposal. The industry has progressed to an understanding that easy answers are not apparent and that evaluation of the environmental stresses associated with any system, even at one moment in time, is a complex undertaking (24). A nonobvious example is the environmental impact of additional employees required for CIP systems operation, such as in their commuting (25).

Finally, consider the environmental impact of each component throughout its entire existence. Life-cycle analysis (LCA) examines the aggregate consequences of system components, not just during their use, but from manufacture to disposal (26). All things considered, most conclusions support acceptability for the relative ecological consequences of SUS (27).

QbD and PAT
Concurrent with SUS, the industry is adopting such initiatives as quality by design (QbD) and process analytical technology (PAT). It turns out that many features of SUS actively support both: e.g., modularity, reconfigurability, flexibility in process flow and layout, and improved compliance values. Whether SUS are applied at process development or full manufacturing scale, their features support these new regulatory initiatives through development of increased process understanding and improved process life-cycle management (3).

Flexibility
Support of so-called flexible manufacturing under CGMP compliance has emerged as a hallmark of SUS-based manufacturing. Economic, regulatory, and market pressures create a demand to include “flexibilities” in biomanufacturing process design: e.g., efficient accommodation of changes in campaign schedules, production scale, and system reconfiguration in support of multiple product portfolios — even to support the active use of multiple production platforms.

Although such power is remarkable in itself, the concept of flexible manufacturing also confers the efficient support over a product’s life cycle of improvements in certain unit operations and even the replication or relocation of entire facilities.

Many flexible features of disposables support even further capabilities or economy. For example, the ease with which a manufacturing process based on SUS can be replicated at a second location facilitates the use of a CMO for production at early stages or in high-risk ventures, allowing the exact process to be readily transferred to a sponsor’s facility when the product reaches a more advanced stage. The promise of such capabilities has not only been demonstrated in a number of real-life SUS implementations, but also has become a major theme in modern biomanufacturing process design (9, 28).

Performance
Many SUS have by now demonstrated the performance they initially promised (Figure 2). Although most disposable mixers and bioreactors on the market are distinct engineering solutions, for example, the performance of many have been established through years of application. Wave-style bioreactors have been used in actual API production seed trains for over five years. Newer stirred-tank SUS reactors have demonstrated excellent performance through peer-reviewed case studies and conference presentations (22, 29). Surprisingly, it is not uncommon to find performance of disposables to be even superior to classical systems (30). The era of single-use-based biomanufacturing has arrived, and many more advanced systems are currently used to manufacture products now in clinical trials.

References
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Table 3: Many demands on single-use systems (SUS) have been met, but many remain for certain unit operations and available systems.

<table>
<thead>
<tr>
<th>Current Concern</th>
<th>Progress</th>
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<tr>
<td>Scalability limitations: 2,000-L working volumes currently the largest available</td>
<td>Higher yields, personalized medicine, and so on are reducing required process volumes.</td>
</tr>
<tr>
<td>Processes and models in infancy: based on preliminary data</td>
<td>Many publications, filings, and case studies are appearing.</td>
</tr>
<tr>
<td>Operational &quot;robustness&quot; unproven: related to above issues</td>
<td>For some SUS, multiple years’ worth of success is being published.</td>
</tr>
<tr>
<td>Disposable materials storage requirements</td>
<td>Process footprint is reduced.</td>
</tr>
<tr>
<td>Questions regarding product and process materials compatibility</td>
<td>Compatibility is being established through vendors, BPSA, and contract studies.</td>
</tr>
<tr>
<td>Inventory, storage, and repetitive purchases required</td>
<td>If anticipated, these are compensated for by reduced operational demands/footprint.</td>
</tr>
<tr>
<td>Potential for vendor dependency (single-source equipment)</td>
<td>For many disposables, multiple vendors have emerged.</td>
</tr>
<tr>
<td>Potential for dependency on single or novel systems</td>
<td>For many disposables, multiple related systems have emerged.</td>
</tr>
<tr>
<td>Observed variability in vendor &quot;maturity&quot; and capacities</td>
<td>Most vendors are advancing as SUS use continues.</td>
</tr>
<tr>
<td>New and different vendor audit issues</td>
<td>Understanding and planning are in progress.</td>
</tr>
<tr>
<td>Availability: generic, standard, or specialized</td>
<td>Less of an issue as this becoming understood</td>
</tr>
<tr>
<td>Novel validation demands</td>
<td>Commercial vendors, BPSA, and contracted studies are establishing methods.</td>
</tr>
<tr>
<td>Novel engineering and process layout/flow design activity</td>
<td>Issue is being addressed (e.g., through a number of experienced consultants).</td>
</tr>
<tr>
<td>Developing industry and regulatory standards</td>
<td>Standards are in progress through aggressive pharma, FDA, and BPSA activity.</td>
</tr>
<tr>
<td>Product containment failure concerns</td>
<td>Previous failure points are being reengineered.</td>
</tr>
<tr>
<td>Potential for contents transfer flow rate limitations</td>
<td>Advanced designs (e.g., larger ports, improved connectors) are resolving this issue.</td>
</tr>
<tr>
<td>Connectivity issues: standardization and continued solutions desired</td>
<td>Numerous connection solutions are appearing in the market.</td>
</tr>
<tr>
<td>Questions regarding new materials: leachables and extractables</td>
<td>Questions are being answered by vendors, BPSA, and contracted studies.</td>
</tr>
<tr>
<td>Questions regarding the carbon footprint and fate of used materials</td>
<td>Individual issues remain, but overall SUS equal to or less than traditional systems.</td>
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</table>


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