Development and Application of Flow-Cytometric Techniques for Analyzing and Sorting Endospore-Forming Clostridia†‡

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The study of microbial heterogeneity at the single-cell level is a rapidly growing area of research in microbiology and biotechnology due to its significance in pathogenesis, environmental biology, and industrial biotechnologies. However, the tools available for efficiently and precisely probing such heterogeneity are limited for most bacteria. Here we describe the development and application of flow-cytometric (FC) and fluorescence-assisted cell-sorting techniques for the study of endospore-forming bacteria. We show that by combining FC light scattering (LS) with nucleic acid staining, we can discriminate, quantify, and enrich all sporulation-associated morphologies exhibited by the endospore-forming anaerobe Clostridium acetobutylicum. Using FC LS analysis, we quantitatively show that clostridial cultures commonly perform multiple rounds of sporulation and that sporulation is induced earlier by the overexpression of Spo0A, the master regulator of endospore formation. To further demonstrate the power of our approach, we employed FC LS analysis to generate compelling evidence to challenge the long-accepted view in the field that the clostridial cell form is the solvent-forming phenotype.

Microbiology traditionally focused on analyzing microbes at the population scale, limiting conclusions about individual cell behavior to culture-level data (4). However, it is well accepted that microbial cultures, even clonal populations, exhibit more cellular heterogeneity than previously appreciated (1, 2, 6, 12, 13, 35). The study of culture heterogeneity is a relatively new field of microbiological research very significant to endospore-forming bacteria, notably clostridia.

Clostridia are obligate anaerobe, gram-positive, sporulating firmicutes that include pathogenic species significant to human/animal health and physiology and non-pathogenic species relevant to the commercial conversion of renewable resources into biofuels and commodity chemicals (26). Pathogenic species such as Clostridium botulinum, C. difficile, C. perfringens, and C. tetani produce 18% of all known bacterial toxins, making it the “most toxic” bacterial genus (28). Cellulolytic and solventogenic species such as C. thermocellum, C. saccharobutylicum, C. cellulolyticum, and C. acetobutylicum are some of the best-studied biomass-metabolizing bacteria, with significant potential for sustainable biofuel production via consolidated bioprocessing (23, 24). Nontoxicogenic, proteolytic species, namely, C. sporogenes and C. novyi, are being engineered into promising chemotherapeutic vehicles, a process called clostridium-directed enzyme prodrug therapy (CDEPT) (25). CDEPT entails the intravenous delivery of clostridial spores, which infiltrate and selectively germinate in the hypoxic regions of solid tumors. The vegetative clostridia then produce an enzyme that acts upon a prodrug precursor, converting it into its toxic form (25).

Central to all these fields of research is sporulation, which occurs in all clostridia in response to still largely unknown signals (26). Sporulation is directly linked genetically to CPE enterotoxin and C2 toxin expression in C. perfringens and C. botulinum (11), respectively. Sporulation is the very reason why C. sporogenes and C. novyi can be administered systemically for CDEPT, and sporulation is believed to be genetically coupled to the solvent metabolism of industrial organisms such as C. acetobutylicum (19). Although significant advances are being made (19), clostridial sporulation remains largely uncharacterized in part due to culture heterogeneity. Only a small portion of a typical culture sporulates for many of the most-studied species (14, 22, 27, 43). Thus, genetic, proteomic, and metabolite analyses are often complicated by culture heterogeneity.

Prior efforts at single-cell analysis of clostridial cultures include Fourier transform infrared spectroscopy, Raman spectroscopy coupled with microscopy, and electro-optical measurements (20, 30–33). Although powerful, these techniques are not capable of high-throughput analysis, are less capable of quantifying bulk population heterogeneity, are unable to enrich phenotypes for subsequent analyses, and often require custom-made equipment. Flow cytometry (FC) is a high-throughput, multiparametric analysis technique capable of discriminating and sorting individual cells. Although microbial FC analysis has been advancing for nearly half a century (35), it is employed to a very low extent for microbial compared to mammalian cell culture analysis. Bacteria are approximately 3 orders of magnitude smaller in mass and contain much less DNA, RNA, and protein per cell than mammalian cells (34). Thus, bacterial light scattering (LS) occurs at or near the threshold level of instrumental noise, and fewer fluorescently
labeled molecules per cell complicate fluorescence detection while increasing sample variance (34). Additionally, there are far fewer established assays that can be employed across all classes of bacteria due to the enormous heterogeneity among them. For example, there are many reports of ambiguous cell viability results with the increasingly popular Live/Dead BacLight viability assay from Invitrogen (3, 17, 36). Furthermore, many interesting bacterial species, including all clostridia, are difficult to genetically modify. Lastly, these are obligate anaerobes, and thus fluorescent proteins like green fluorescent protein cannot be employed due to strict requirements for oxygen (39).

Here we report the development of FC and fluorescence-assisted cell-sorting (FACS) methods for investigating clostridial sporulation and culture heterogeneity. We demonstrated the superior multiparametric detection, high-throughput, and sorting advantages of these techniques. Discriminated, quantified, and enriched all sporulation-associated morphologies, discussed in detail elsewhere (19). We validated our assays by analyzing and sorting batch cultures of wild-type (WT) C. acetobutylicum ATCC 824 and mutant or recombinant strains (8, 15, 40). We sorted all sporulation-associated cellular morphologies for microscopic analyses and developed a novel assay for determining the metabolic state of clostridial cultures. Finally, we demonstrated the utility of our methods by correlating FC-detected morphology changes with bulk-phase metabolite concentrations and generated compelling evidence to challenge the long-accepted view in the field that clostridial-form cells are responsible for solventogenesis (18).

MATERIALS AND METHODS

Bacterial strains and plasmids. Bacterial strains employed in this study were C. acetobutylicum ATCC 824 (ATCC, Manassas, VA) and the degenerate strain M5 (which has lost the 192-kb, 176-gene megaplasmid pSOL1) (5, 8). Plasmids employed were pSOS95del (MLS' [macrolide-lincosamide-streptogramin B resistance], CoElI origin of replication, repl. [pLM13 origin of replication], Amp' [ampicillin resistance], blu' promoter) (40) and the Spo0A expression plasmid pMSPO0A (Amp', MLS', CoElI origin of replication, repl., carries Spoo4) (15).

Growth conditions and maintenance. C. acetobutylicum ATCC 824 and recombinant strains were grown anaerobically in 2 YNTG broth supplemented with 15% glycerol and revived by anaerobic plating on 2 YTG.

Results

Analytical methods. Cell growth was monitored by measuring optical density at 600 nm ([600] of pSOL1. Two hundred milliliter static flask cultures were used for all samples to kill all vegetative cells, to induce spore germination, and to insure the presence of both spores and vegetative cells. Samples were diluted to ~10<sup>6</sup> cells/ml in 5 to 6 ml of 1% NaCl. Sorted cells were cultured in 15-ml Falcon tubes for two-channel sorting and in 10-ml FC tubes for four-channel sorting. The sample injection port and sample collection apparatus were both maintained at 4°C. The final concentration of sorted cells was much lower than the sample concentration—these values were ~10<sup>6</sup> to 10<sup>8</sup> cells/ml versus ~10<sup>6</sup> cells/ml—and total volumes were on average 25 to 50 ml. Subsequently, staining characteristics were lost after sorting due to the low concentration of dyes, as confirmed by microscopy and FC. Sorted populations were pelleted by a 15-min, 25,000 × g centrifugation. Pellets were resuspended in 3 to 10 µl depending on pellet size.

LS sample preparation. Samples for FC analysis were prepared by pelleting 500 µl of culture via a 1-min centrifugation at 16,000 × g. The supernatant was discarded and cells were washed by resuspension in 1 ml of sterile filtered 1% (wt/vol) NaCl. Cells were pelleted again and then resuspended in 40 µl of sterile filtered 1% (wt/vol) NaCl. This is the same protocol for preparing cell samples for nucleic acid (NA) staining analysis. The LS sample was prepared by diluting 1 to 5 µl of washed cells in 1 ml of sterile filtered 1% (wt/vol) NaCl.

NA staining protocol. For an A<sub>600</sub> of <0.5, 1 ml of culture was sampled. For an optical density at 600 nm of >0.5, 0.5 ml of culture was sampled. Cells were washed and resuspended as described for LS samples. The 40 µl of washed cells was diluted to a final volume of 200 µl with sterile filtered 1% (wt/vol) NaCl. The Invitrogen Live/Dead BacLight bacterial viability stain was employed. Syto9 and PI were mixed to final solution concentrations of 1.67 mM and 9.985 mM, respectively. Two microliters of Syto9-PI mix was added to the 200-µl sample, and this mix was used for sorting. Samples were fixed, prepared, and analyzed as previously described (19).

Transmission electron microscopy (TEM). Samples were fixed, prepared, and imaged as previously described (19).

Metabolic flux analysis. Metabolic flux calculations were performed using a previously developed model of C. acetobutylicum primary metabolism (10). Recent developments using a genetic algorithm improved model convergence in the presence of the nonlinearities (R. S. Senger and E. T. Papoutsakis, unpublished data).

RESULTS

LS FC analysis of sporulating cells [824(pSOS95del) and WT strains] identifies reproducible temporal patterns distinct from those of the asporogenous strain M5. (i) Analysis of sporulating strain 824(pSOS95del). We focused our initial investigations on cultures of 824(pSOS95del), which is an expression plasmid control strain and has been shown to sporulate in large proportion of the culture compared to what is seen for the WT ATCC 824 strain (15). This phenomenon is shown to occur for all tested plasmid control strains due to host-plasmid interactions (38, 41). We examined the LS characteristics of three 824(pSOS95del) batch cultures (Fig. 1A and B), capturing the full sporulation program and metabolism
FIG. 1. LS investigation and metabolite analysis of a typical sporulating 824(pSOS95del) static flask culture. (A) Metabolite profiles, growth curves, and identification of batch culture stages. Symbols: ×, acetate; ▲, butyrate; □, acetone; Δ, ethanol; and ○, butanol. Batch culture stages are identified by black bars as follows: 1, exponential; 2, transition; 3, early stationary; 4, mid-stationary; and 5, late stationary. (B) Dot plots from the LS (FSC/SSC) analysis. Dot plots were gated into four regions as follows: A, low FSC/high SSC; B, mid-FSC/high SSC; C, high FSC/high SSC; and D, all FSC/low SSC. (C) Composition of cell population for each LS region in the panel B dot plots shown as the percentage of total population. (D) Phase-contrast microscopy of the same sample analyzed by FC. Numbers on microscopy images correspond to the numbers on dot plots. Arrows point to the morphologies we are most interested in identifying and quantifying by FC. conc., concentration.
of \textit{C. acetobutylicum} ATCC 824. LS samples were taken every 2 h during the exponential and transition phases, every 4 h during early stationary phase, every 6 to 12 h during mid-stationary phase, and every 12 to 24 h during the late stationary phase (Fig. 1A). FC analyses of three biological 824(pSOS95del) replicates revealed four unique cell populations, which are gated as follows: A, low-intensity FSC (low FSC)/high-intensity SSC (high SSC); B, middle-intensity FSC (mid-FSC)/high SSC; C, high FSC/high SSC; and D, all FSC/low SSC. FSC/SSC dot plots and corresponding phase-contrast microscopy from one representative 824(pSOS95del) culture are shown in Fig. 1B and D. Populations A and D were the two major populations during exponential growth, comprising 69% and 29% of the total population, respectively. Phase-contrast microscopy demonstrated that all cells were rod shaped and presumably vegetative. During the transition phase, the majority of cells (>91% of the total population) exhibited greater SSC intensity, condensing into population A. The majority of cells still appeared rod shaped and vegetative, but there was a very small emerging population of swollen, dark-phase cells, characteristic of the cigar-shaped, dark-phase clostridial-form cells (18, 22). As the culture progressed into early stationary phase, 23% of the population exhibited higher FSC intensity, as illustrated by the evolving population in gate B. Phase-contrast microscopy revealed the presence of clostridial-form cells (swollen, typically bright-phase morphologies) and forespore-containing cells (swollen morphologies, exhibiting a single, bright-phase end) in approximately one-fourth of the population. During mid-stationary phase, a population with FSC of even greater average intensity emerged, illustrated by the cells accumulating in gate C. The total cell population was distributed among gates A, B, and C as 61%, 21%, and 18% at this culture phase, respectively. Phase-contrast microscopy revealed the presence of rod-shaped cells, clostridial-form cells, forespore-containing cells, endospore-containing cells (bright-phase spores still surrounded by the mother cell), and free spores (very-bright-phase spheres). During late stationary phase, the majority of the population resolved in gates A and C, comprising 73% and 19% of the total population, respectively. A small percentage of cells exhibited gate B LS characteristics during late stationary phase (7.9%) compared to those exhibiting them during mid-stationary phase (20.7%). Phase-contrast microscopy revealed free spores, endospore-containing cells, and rod-shaped cells but few clostridial-form cells and forespore-containing cells compared to what was seen for early- and mid-stationary-phase samples. LS population percentages for each phase of growth are summarized in Fig. 1C.

This and biological replicate analyses suggest that the increasing mean FSC populations in gates B and C are representative of sporulating cells, more specifically, clostridial-form cells and forespore-containing cells in gate B and endospore-containing cells and free spores in gate C. To further substantiate our claim, we applied the same analysis to WT and M5 batch cultures.

(ii) Analysis of the sporulating WT and asporogenous M5 strains. For WT cultures, we expected the same LS characteristics as for of 824(pSOS95del) but as a smaller proportion of the total cell population (15). M5 does not sporulate and does not produce clostridial-form cells, and thus we do not expect to observe well-defined populations in gates B and C. Three biological replicates of WT and M5 strains were analyzed for LS characteristics as for the 824(pSOS95del) cultures. Dot plot samples are shown in Fig. S1A and B in the supplemental material. The same LS characteristics and temporal phenomena observed for 824(pSOS95del) cultures were apparent for WT cultures. Again, there was the condensation of cells into gate A, followed by the evolution of populations in gate B and then gate C, and finally the resolution of distinct populations in gates A and C that were not connected by a significant gate B population (Fig. 1B, “Late Stationary” panel). However, as expected (15), a smaller percentage of the total population was involved. For any WT sample, there were at most 14.7% and 8.8% of the total population measured in gates B and C, respectively, compared to 22.8% and 18.8% in gates B and C for a typical 824(pSOS95del) culture.

M5 cultures never displayed distinct populations in gates B and C at any time. They displayed an increase mean SSC intensity during the switch from exponential to transition phases, but beyond the transition phase, the cells either retained gate A LS characteristics or exhibited disperse and increased FSC/SSC intensities (Fig. 2). Interestingly, the observation of disperse, increased FSC/SSC intensity coincided with the end of metabolism, marked by the cessation in glucose consumption and metabolite production (Fig. 2). This was exhibited by all M5 cultures and other mutant strain cultures that prematurely ceased metabolism (data not shown).

NA (PI and Syto9) staining in combination with LS allows discrimination of all major sporulation-associated phenotypes, and FACS makes it possible to isolate highly enriched sporulation phenotypes. So far, our data have shown that LS alone is capable of distinguishing and quantifying sporulating from nonsporulating cells but that it does not clearly discriminate between the clostridial-form cells and forespore-containing cells (presumably gate B) or between endospore-containing cells and free spores (gate C). To solve this problem, we combined the LS assay with the Live/Dead BacLight viability assay. Although the BacLight assay’s intended application is to distinguish between live and dead bacteria, several investigators have described ambiguous results in using this assay as a viability test (3, 17, 29, 36). In the context of clostridia or other endospore formers, the Live/Dead BacLight assay was used for analyzing vegetative \textit{C. acetobutylicum} cultures via fluorescence microscopy, but results for sporulating phenotypes were not reported (16). We recently argued (19) that this assay cannot be used as a viability assay in this organism, but it can likely distinguish sporulation phenotypes.

We analyzed the same cell samples from the previously described 824(pSOS95del), WT, and M5 cultures. There were no obvious correlations between PI and Syto9 NA staining alone and sporulation phenotypes, but when gated upon the LS gates, we noticed distinct subpopulations in LS gates A, B, and C. As shown in Fig. 3A, two distinct NA staining populations resolved for the early-stationary-phase sample, which we gated as R1 (PI+/Syto9+) and R2 (PI+/Syto9dim). For the total population, 34.3% and 43.8% of the cells were in gates R1 and R2, respectively. When gated upon LS region A, 27.4% and 48.3% of the cells were in gates R1 and R2, respectively. When gated upon LS region B, 50.1% and 33.5% of the cells were in gates R1 and R2, respectively. Thus, a noticeable enrichment for R1
(PI−/Syto9+) was witnessed in LS region B and likewise for R2 (PI−/Syto9dim) in LS region A.

To more rigorously characterize these subpopulations, we employed FACS to sort and analyze cells via phase-contrast microscopy and TEM. We sorted at least 5 million cells from all four populations: R1, A; R2, A; R1, B; and R2, B. We found that 5 million events are necessary to obtain enough cells for efficient pelleting via high-speed centrifugation. As shown in Fig. 3B, there was enrichment for clostridial-form cells in R1, B; for forespore-containing cells in R1, A; and for rod-shaped cells in both R2, B and R2, A, presumably vegetative cells. The purity for each specific phenotype was never 100%, because sorting was performed at rates of >80,000 cells/s. Even at these rates, it still required ca. 5 h per sample to obtain 5 million cells for each population. Thus, for practical considerations, we sacrificed purity for large numbers of cells.

Enrichment for clostridial-form cells in LS gate B was expected, but enrichment for forespore-containing cells in LS gate A was not. Postsorting LS analysis of the enriched populations revealed the following mean FSC intensities: 2,028 for R1, B (clostridial-form cells); 1,735 for R2, B (rod-shaped cells); 1,250 for R1, A (forespore-containing cells); and 1,037 for R2, A (rod-shaped cells). Gate B FSC bounds are 1,450 to 3,500, suggesting that clostridial-form cells and a small population of rod-shaped cells are captured in gate B. The forespore-containing cell population exists at the boundary between gates A and B, which we effectively distinguish from the bulk of rod-shaped, lower-FSC-intensity gate A cells via NA discrimination. However, to definitively capture forespore-containing cells and clostridial-form cells from the bulk of rod-shaped cells via LS discrimination alone, we extended the FSC lower bound of gate B to 1,200, sorted from the rest of the population, and imaged via TEM. Sorting for TEM analysis was performed at <15,000 cells/s to increase sample purity. Postsorting LS analysis revealed that the LS gate B population was enriched to 64% of the total population from 27% in the presorted population, and TEM imaging revealed that both clostridial-form cells (Fig. 4A) and forespore-containing cells (Fig. 4B) were the predominant morphologies.

In order to discriminate between endospore-containing cells...

FIG. 2. LS characteristics of an asporogeneous M5 culture. (A) LS dot plots from various stages of the batch culture. The broken gate encompasses the disperse high-intensity FSC-SSC events, which are characteristic of a metabolically inactive culture and absent from an actively metabolizing culture. (B) Metabolite profiles and growth curve for a typical M5 batch culture. Symbols: ○, glucose; ▲, acetate; ●, butyrate; and △, ethanol. There was no measurable butanol or acetone. Samples corresponding to the dot plots in panel A are marked with numbers as follows: 1, exponential; 2, transition; 3, mid-stationary; and 4, late stationary. Notice that between hours 20 and 25, metabolite concentrations do not change appreciably, suggesting that the culture is no longer metabolically active. This corresponds to the time at which disperse high-intensity FSC-SSC events become more prevalent and the culture is unable to be serially cultured. conc., concentration.
FIG. 3. Refinement of LS investigation with NA staining for both early- and late-stationary-phase samples of an 824(pSOS95del) batch culture. Populations were sorted based upon the combination of LS and NA staining characteristics. (A) LS and NA staining dot plots for an early-stationary-phase sample. We sorted the following four populations by FACS: R1, B (PI* /Syto* and mid-FSC/high SSC); R2, B (PI*/Syto** and mid-FSC/high SSC); R1, A (PI*/Syto* and low FSC/high SSC); and R2, A (PI*/Syto** and low FSC/high SSC). Percentages within each NA gate for the respective LS gate are indicated. (B) Microscopy of sorted populations from the early-stationary-phase sample. (C) LS and NA staining dot plots for the late-stationary-phase sample. The following two populations were sorted by FACS: R3, C (Syto*/PI dim and high FSC/high SSC); and R4, C (Syto**/PI dim and high FSC/high SSC). The percentage within each NA gate for the respective LS gate C is indicated. (D) Microscopy of sorted populations from the late-stationary-phase sample. (E) Summary of the predominant morphologies for each sorted population. Postsorting analysis revealed that forespore-containing cells exhibited LS characteristics very near the border of gates A and B. We expanded gate B and sorted again to show that gate B captures predominately clostridial-form cells and forespores (Fig. 4).
and free spores, we performed NA staining on late-stationary-phase 824(pSOS95del) samples. Multiple NA staining populations resolved (Fig. 3C, top right), but upon gating upon LS region C, only two distinct populations remained (Fig. 3D, bottom right). We gated these as R3 (PIdim/Syto9\(^+\)) and R4 (PIdim/Syto9\(^{dim}\)), sorted, and imaged via phase-contrast microscopy. Phase-contrast microscopy revealed that 75% of R3, C-sorted cells were endospore-containing cells or free spores (top two images of Fig. 3D), and postsorting LS analysis revealed that 88% of sorted cells exhibited gate C LS characteristics. Significantly, microscopy analysis showed that 90% of the spore morphologies were endospore-containing cells.

Phase-contrast microscopy of R4, C-sorted cells suggested that 50% of the cells display a spore morphology, of which >95% were free spores (bottom two images of Fig. 3D). FC postsorting analysis of R4, C cells revealed that 27% exhibited gate A, 5% gate B, and 68% gate C LS characteristics. The discrepancy between FC and microscopy postsorting analysis was likely due to poor microscope slide preparation, with many cells being out of the plane of focus. Thus, we sorted upon LS gate C characteristics only and analyzed by TEM to more rigorously characterize the sorted morphologies. Sorting was performed at <15,000 cells/s in order to increase sorted population purity. Postsorting analysis revealed that 91% of the cells exhibited LS gate C characteristics. Inspection of TEM images from LS gate C-sorted cells revealed that >90% of the morphologies were of either endospore-containing cells or free spores (Fig. 4C and D).

**FC analysis provides precise temporal insight into sporulation and identifies multiple rounds of sporulation and spore germination in a single batch culture.** We applied LS analysis to quantitatively assess sporulation kinetics. We first analyzed and compared batch cultures of 824(pMSPO0A), which is a Spo0A overexpression strain under its natural promoter (copy number, ca. 7), against those of the WT and 824(pSOS95del) strains. Spo0A is the master transcription regulator of both sporulation and solvent formation in clostridia, and upon overexpression during vegetative growth it is suggested to induce sporulation earlier than what is seen for both WT and 824(pSOS95del) cultures (15). In order to determine and accurately compare when sporulation begins, we accounted for differences in inocula and the lengths of lag phase by setting time zero (\(t_0\)) as the time at which the batch culture reached an \(A_{600}\) value of 1.0. Distinct and defined populations in gates B and C appeared 9 hours and 15 hours early in the 824(pMSPO0A) culture compared to the average of three biological replicate 824(pSOS95del) and WT cultures, respectively (see Fig. S2 in the supplemental material). Our data also show that the 824(pSOS95del) initiates sporulation 6 hours earlier than WT and that for all cultures it takes approximately 10 to 14 h for clostridial-form cells to differentiate into endospore-containing cells.

LS analysis of these cultures also showed that batch C. acetobutylicum cultures can and often do carry out multiple rounds of sporulation. As shown in Fig. 5, top three graphs, the distinct gate C population (endospore-containing cells and free
spores) essentially disappeared from the WT culture between hours 85 and 95 and then reappeared 15 to 25 h later as a greater percentage of the total population. The same phenomena occurred for 824(pSOS95del) cultures. As shown in Fig. 5, bottom three graphs, the percentage of cells exhibiting gate C characteristics was reduced from 18% to 6% of the total population between hours 52 and 60 and then reappeared as almost 50% of the total population 20 to 30 h later. These results suggest spore germination and multiple rounds of sporulation, which was previously suggested but not proven (38), and is in accordance with previous DNA microarray analyses (19). Additionally, the abrupt drop in spore population suggests synchronized germination. Multiple rounds of sporulation occurred in all but one of the sporulating batch cultures that we analyzed during this study (seven of eight total), and the phenotypes seen exhibited interesting correlations to culture metabolism, which we discuss next.

Correlation of LS-discriminated phenotypes to cellular metabolism: is the clostridial cell form responsible for solvent formation? Using C. saccharobutylicum, previously classified as C. acetobutylicum P262 (21), Jones et al. suggested from microscopic evidence that a correlation exists between the number of swollen, cigar-shaped clostridial-form cells and solvent production (18). Using chemical mutagenesis, they derived mutants unable to form clostridial-form cells, which did not produce solvents. Other mutants that displayed reduced glucose accumulation produced intermediate levels of solvents, and mutants blocked at the clostridial-form cell phenotype produced WT levels of solvents. They concluded that clostridial-form cells were responsible for the conversion of acids to...
solvents, which is now widely accepted for all solventogenic clostridia. Our data challenge this view for C. acetobutylicum ATCC 824.

By comparing butanol fluxes to LS populations, our data showed that the fraction of the vegetative, rod-shaped population is directly correlated to butanol flux; in contrast, the frequency of clostridial-form cells and of endospore-containing cells plus free spores is inversely correlated to butanol flux (Fig. 6). We determined butanol flux as the change in butanol concentration over time per unit biomass (in $A_{600}$ units, as determined by optical density) (10). The fraction of vegetative cells and the butanol flux increased and decreased largely in synchrony. In contrast, the fraction of clostridial-form cells followed a largely opposite pattern in terms of butanol flux correlation through hour 50. The population of clostridial-form cells remained relatively constant from hour 50 on, while the fraction of the endospore-containing cell/free spore population continuously increased from hour 60 on. At hour 24, the endospore-containing cell/free spore population began to increase with an approximately 10- to 14-h lag behind the clostridial-form cell population and at the expense of clostridial-form cell population. The endospore-containing cell/free spore population reached a maximum for the first round of sporulation around hour 45 to hour 50 and then decreased, presumably due to germination. The germination process persisted for 10 to 15 h and was accompanied by increases in the percentage of vegetative cells and in butanol flux. A biological replicate 824(pSOS95del) batch culture exhibited the same phenomena, but with a less dramatic germination event and subsequently less of an impact on butanol flux (data not shown). Similar phenomena were observed for WT cultures (see Fig. S3 in the supplemental material) but were not as pronounced, since a smaller percentage of the culture sporulates. These data suggest that a vegetative cell phenotype is predominately responsible for butanol production, and such a vegetative cell phenotype is likely a precursor to clostridial-form cells.

**DISCUSSION**

We developed FC methods and demonstrated the ability to analyze C. acetobutylicum cultures faster, more precisely, and more quantitatively than ever before. By LS discrimination alone, we were able to detect the temporal aspects of sporulation, accurately quantify the proportion of the population participating in sporulation, and sort cultures into enriched populations for subsequent analysis. By coupling with NA staining, we effectively discriminated between all four major sporulation-associated phenotypes, and by using FACS, we were able to enrich for the various sporulation phenotypes. Finally, we demonstrated that previously supposed phenomena and new hypotheses could be more critically examined using FC. For example, using the FC LS assay, we detected and characterized multiple rounds of sporulation along with germination and early sporulation due to the overexpression of the Spo0A. Both phenomena were previously proposed (15, 38) but never proven or measured with such precision. We also determined that disperse high-intensity FSC-SSC events are characteristic of metabolically inactive cultures. We suggest that these events are due to doublets and triplets of lysed cells, which could exhibit a stronger propensity to aggregate than nonlysed cells.

In addition to refining the LS assay, NA staining combined with LS revealed important physical and biochemical information about sporulation phenotypes and will make possible more-critical analysis of such phenotypes via FACS enrichment in the future. For example, the exclusion of all NA stains from the free spore but not the endospore in endospore-containing cells suggests an addition to the developing spore, likely the spore coat, which is responsible for Syto9 exclusion. Interestingly, our observation of Syto9 and PI exclusion by free spores and partial exclusion by endospores is similar to that reported for the analysis of Bacillus cereus endospore germination via Syto9 staining (9) and identical to results for Syto13 and PI staining of Paenibacillus polymyxa (7). Also, the exclusion of PI from clostridial-form cells and forespore-containing cells but not from the majority of rod-shaped cells and the mother cell compartments of the endospore-containing cells suggests a physical difference in the cell membranes of these phenotypes. Cell membrane composition changes are a well-known fact in C. acetobutylicum and other solventogenic clostridia. Particularly during the metabolic shift from acidogenesis to solventogenesis, cells compensate for the increase of membrane fluidity associated with increased butanol concentration by increasing the proportion of saturated fatty acids and the mean acyl chain length in the cell membrane (37, 44). However, this knowledge comes from heterogeneous cultures, and little is known about specific phenotypes. We demonstrated that FACS can be employed to rapidly purify and quantify all sporulation populations for subsequent detailed TEM analysis and should be able to complement other single-cell analyses, such as Raman and Fourier transform infrared spectroscopy (31–33), which precisely measure membrane composition, DNA content, and RNA content, etc.

To conclude our discussion of method development, we applied these techniques to investigate the relationship between morphology and solvent production. We showed that correlations between LS analyses and metabolite production for both WT and 824(pSOS95del) batch cultures strongly suggest that the clostridial-form cells do not produce solvent and actually detract from solvent production. Subsequently, we proposed that a clostridial-form cell precursor is actually the major solvent producer, which challenges a long-accepted belief that clostridial-form cells are the major solvent producer in all solventogenic clostridial cultures.

Although we report only on the analysis of four C. acetobutylicum strains, we applied our methods to several other recombinant strains and do not foresee why such methods cannot be immediately applied to all clostridial species. However, extra care should be taken in the analysis of pathogenic species such as C. botulinum, so that no aerosolized toxin is generated during FC analysis. Given the tremendous interest in clostridial cell culture, we look forward to the application of FC techniques for accelerating research on various industrially significant clostridial species (C. acetobutylicum, C. thermocellum, C. cellulolyticum, C. phytofermentans, and C. beijerinckii, etc.) and for advancing the study of pathogenic species (C. botulinum, C. difficile, C. perfringens, and C. tetani, etc.).
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