Introduction to DSP – Harvesting

| Your Objectives: | |
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| At the end of the lesson, you sho | ould be able to sequence the steps of harvesting. |
| Harvesting | |
| Cell | is a critical step in connecting upstream monoclonal antibody |
| production with downstream | purification. Selecting the cell |
| · · | on the characteristics of the cell culture process can be de early on in process development; a sound understanding of |
| the current process, as well as | of the and cons of the various cell |
| harvesting | available, is required. |
| Companies will take decisions a | as to which harvesting technique best suits their own process, |
| and as to which variables would | be preferential for . |
| being harvested, and (3) proper | d and equipment depends on (1) the type of cells, (2) product erties of the process fluids. Traditional (standard) techniques tion, tangential-flow filtration (TFF, or crossflow filtration), |
| , an | d depth filtration, as well as specialized solutions that are to be |
| coupled with either microfiltration | on or centrifugation with TFF or then with depth filtration. |
| Centrifugation | |
| Disc-stack centrifugation (e.g. fr | om Alfa Laval Inc. or Westfalia Separator Group Ltd.) has been |
| in use for | harvesting for some time. A disc-stack centrifuge uses |

| , ii | nclined conical discs | to separate out v | vhole cells and large cell |
|--|------------------------|----------------------|----------------------------|
| debris. | | | |
| Shearing, however, can | | cells and thereby | increase the number of |
| submicron particles, particles techniques coupled with secon | | | • |
| filtration can | | further cl | arification, removing |
| sol | id particulates. | | |
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| Filtration | | | |
| The bioprocess industry adop | ted membrane clarifi | cation (purification | on) methods from other |
| technologies; but it has also | | modules | specifically designed for |
| biopharmaceutical purposes, in | cluding for primary cl | arification of fern | nentation and cell-culture |
| systems. When a membrane | technique is | | , the size and fouling |
| potential of the membrane is considered. | | | |
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| Microfiltration is | sensiti | ve to changes in | feedstock quality such as |
| cell culture viability, cell dens | ity, and medium com | ponents. High co | ell densities and low cell |
| viabilities can result in high | transmembrane pre | ssure (|) for |
| constant-flux membranes, the relatively lower for typical | • | | |
| of | feed solution acros | s the membran | e reduces concentration |
| polarization and creates a | pressure drop, fo | rcing a size-exc | clusion condition. Such |
| are | e currently available | with various pore | e sizes, typically between |
| 0.1μm and 1μm for primary ha from Pall Corporation). | arvest (e.g. EMD Milli | oore's ProStak sy | stem and PallSep Biotech |

| Depth filtration retains particles, both larger and smaller | their pore |
|---|-------------------------------------|
| size, throughout their porous media. Particle retention involve adsorption through hydrophobic and ionic (as well as other) int | |
| filtration provides some benefits, b | out is not without its limitations. |
| Commercially available technologies include filter sheets (Seitz Corporation); encapsulated modules (<i>Supracap</i> 200, by Pall Inc. formats (e.g. <i>Millistak+</i> pod depth filter media by Emanuel Mer | .); and disposable, scalable |
| Cell separation using depth filters | |
| Benefits | |
| Low material and hardware costs | |
| Ease of use | |
| Fast process | |
| High flexibility from scale-up options (e.g. for multiprod | uct facilities) |
| Filtration with minimal shearing forces so as to offer cel | l and product protection |
| approved materials | |
| High-quality of filtrate | |
| Space-savings equipment | |

Disadvantages

• Necessity of monitoring differential pressure

| • | for conducting economic evaluation at all stages? |
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- Necessity of the removal and disposal of filters
- Scale-up limitations like budget limits or infrastructural limits

| Tangential-flow filtrat | ion: Methods for harvestir | ig small-scale | | may |
|-------------------------|---|------------------|--|-----------|
| not work for process | scales of between 10,00 | 0 and 20,000 lit | cres. As Ian Sellick o | bserved, |
| "Handling requiremen | ts for such large | | call for employing la | arger and |
| more continuous ty | pes of processing syste | | me form of TFF". ce it retains particul | |
| other molecules too | large to pass through | its membrane | pores, and are tra | nsported |
| | the tangential flow. | | | |
| J | lude both cassettes (<u>flat)</u> npact TFF cassette device | | | • |
| and screens to form | | flow paths that | t prevent product ch | annelling |
| (which occurs when r | nembrane surface area is | not fully used a | and the feed channe | ls to the |
| retentate). He also | describes enhanced hollo | ow-fibre | | for cell |
| • | ve used a nonbinding poly es provide several benef | • | , , , | |
| including cost-savings, | lower | of cros | s contamination, and | d greater |
| manufacturing flexibili | ty. | | | |

 $Source: \underline{https://bioprocessintl.com/downstream-processing/chromatography/a-decade-of-harvesting-methods-331186/$

<u>Notabene:</u> "e.g." = one of other examples (zum Beispiel) / "i.e." = namely (sprich, nämlich, das heisst)

| Optimisation |
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| have been made to correlate the characteristics of a cell culture |
| (e.g. viability, density) and the efficiency of a harvest process. In one study of a high cell density |
| culture, the of the primary recovery depended on cell density and |
| cell viability. High cell densities and low viabilities result in larger numbers of whole cells and |
| solid such as colloids and cell debris. High titres* typically generate |
| from conditions of high cell densities and low viability. Using turbidity as a |
| for product quality, researchers demonstrated a linear correlation |
| between cell culture viability and clarification efficiency a given |
| centrifugation condition. Their also showed that higher feed rates |
| lowered clarification because of lower centrifuge residence times and that the correlations for |
| the centrifuge were the same as those for the subsequent depth |
| filtration step. |
| The increasing use of single-use technologies in upstream processes may |
| a shift in harvest strategies. With traditional fed-batch culture |
| processes, harvest clarification is usually achieved by centrifugation followed by depth filtration. For processes based entirely on disposables, the disc-stack centrifuge needs to be |
| by filtration alone". |
| |

* **Titre**: (US: titer) = concentration of a solution determined by titration, the minimum volume of a solution needed to reach the end point in a titration; the concentration of an antibody, as determined finding the highest dilution at which

it is still able to cause agglutination of the antigen.

Scale Challenges

| The current trend towards a small-scale batch process will necessitate primary recovery |
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| techniques that can provide quick . A system of glass-fibre filters, |
| coupled with a retentive membrane to provide "a rapid solution that can be implemented with very little preparation," might be used. Glass-fibre and polyethersulfone are inherently low in extractables and have been validated for low |
| levels of endotoxin and other contaminants. They are steam sterilizable and are compatible |
| with gamma irradiation, enabling pressure sterilization with method. |
| TFF and centrifuge–depth filtration are methods for harvesting |
| large-scale batches. Process transfer, however, sometimes requires alternative, streamlined approaches. A <i>Genentech</i> study evaluated various dual-layer, single-stage depth filter media to |
| an alternative to its traditional two-stage depth filtration train to |
| be used at a separate facility. The work involved at various |
| turbidities, in a system that demonstrated increased capacity at all |
| levels without significant plugging of the internal pore structure of the depth filters. Researchers further suggest that "one key to increasing future depth-filter capacity may () be |
| to control particulate clustering () or modulate flow rate to or |
| exploit critical shear thresholds within a depth filter." |
| High-density cell cultures |
| As increased during this past decade, so too did the importance of |
| efficient product recovery methods. With monoclonal titers |

| reaching $\sim\!25$ g/L and the expectation of even greater values in the future, clarification of | of high- |
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| density cell cultures will present unique challenges to | . Ever- |
| increasing amounts of cell debris will need to be removed quickly | |
| prevent bottlenecks in downstream processes. | |
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| Aufgahe Lückenteyt | |

Folgende Wörter bitte in den Lückentext einfüllen. Jedes Wort kommt einmal vor. Bitte Gross- und Kleinbuchstaben beachten.

along, antibody, avoid, best, batches, bioreactors, biomanufacturers, by, cell, centrifugation, damage, Depth, developed, diagonal, Efforts, either, FDA, find, for, filters, flow, harvesting, highly, impurities, marker, membranes, modules, Need, provide, pros, quality, replaced, require, resulting, results, risk, shown, smaller, suitable, stacked, technologies, testing, than, them, to, titres, times, TMP, turnarounds, typical, used,