

Modernizing legacy Vaccine processes

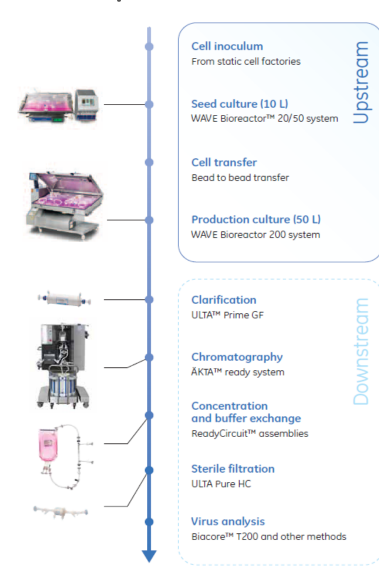
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Imagination at work

Live Influenza virus production



Influenza process overview



Scale-up from small scale to pilot scale in single-use format

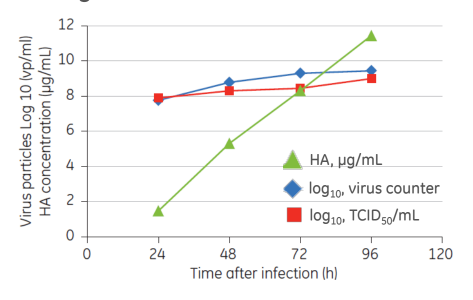
Comparison of culture performance in 10 L and 50 L microcarrier culture in rocking bioreactors

Downstream purification in flowthrough chromatography mode with Capto™ Q and Capto Core 700 chromatography media (resins)



Virus growth kinetics

HA concentration and virus titer during culture



Cell morphology at time of harvest (96 h)



HA concentration at harvest was close to 12 μ g/mL and the virus concentration was > 10 9 infective units/mL



Purification Workflow

NFF

ULTA™ prime **GF**

Microcarrier and cell debris removal Adjustment of conducitvity

Capto Q

Capto™ Q – Flow through
Reduction of DNA and host cell proteins

Capto Core 700 Capto Core 700 – Flow through Reduction of host cell proteins

CFF

ReadyToProcess™ hollow fiber

Concentration, buffer exchange and removal of DNA and host cell proteins

SF

ULTA pure HC

Sterile filtration

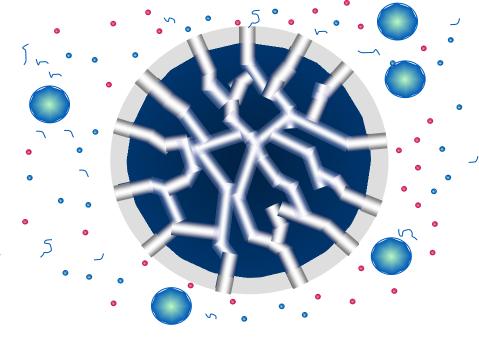


Core bead chromatography, the principle

- Separation of substances with different features such as size, charge, and biological properties.
- The target product may bind or not, the important part is that it does the opposite of what most impurities do!

Highly porous particle, offering a huge surface for binding of proteins

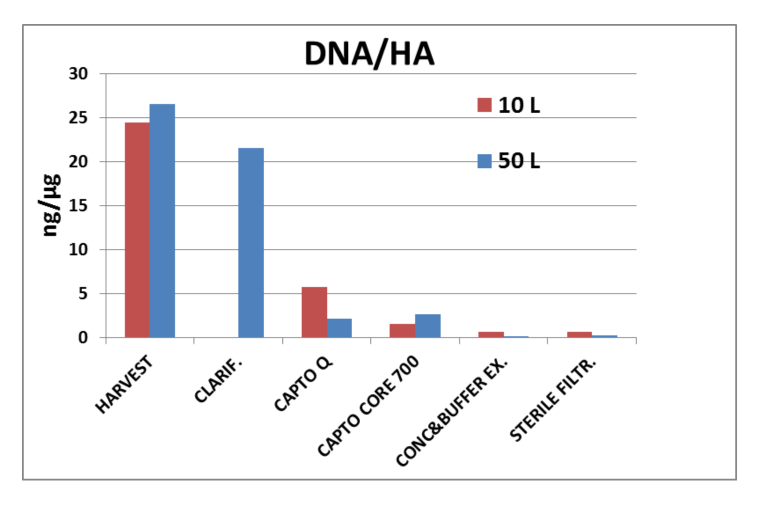
Surface modified to enable selective binding of product or impurities



Spheric particle, 1/10th of a mm in diameter



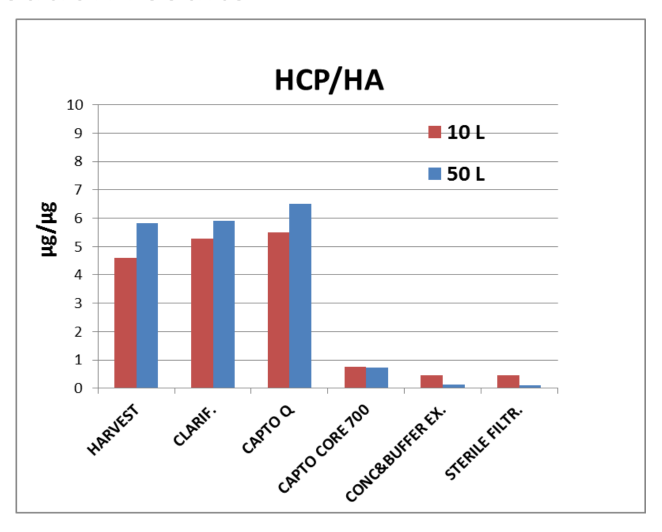
Purification results



Capto™ Q: Reduces host cell DNA



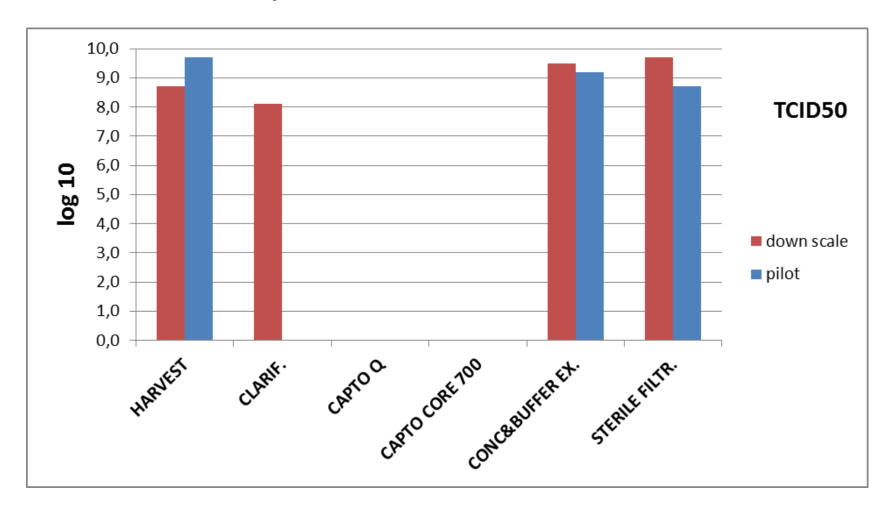
Purification results



Capto™ Core 700: Reduces host cell protein



Virus infectivity



Process does not impair virus infectivity



Yellow fever virus propagation – from eggs to cells



GMP manufacturing of viral vaccine



Xcellerex™ XDR-50 bioreactor

Vero cells (WHO-10-87)

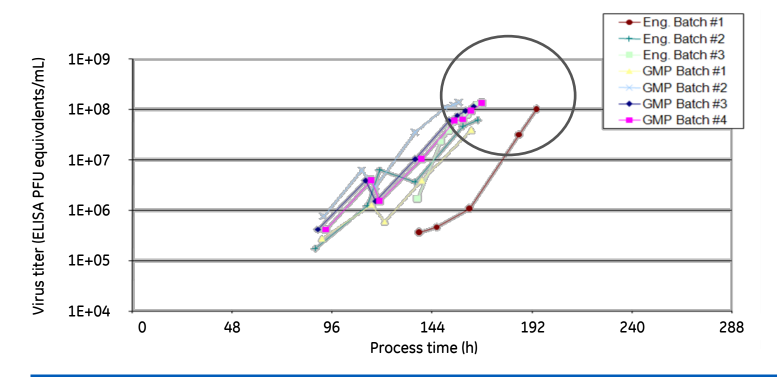
- Cytodex[™] 1 microcarrier
- Serum free, animal componentfree medium

Yellow fever virus 17D



Virus production drain down refeed

PFU equivalents from Eng and GMP bioreactor runs

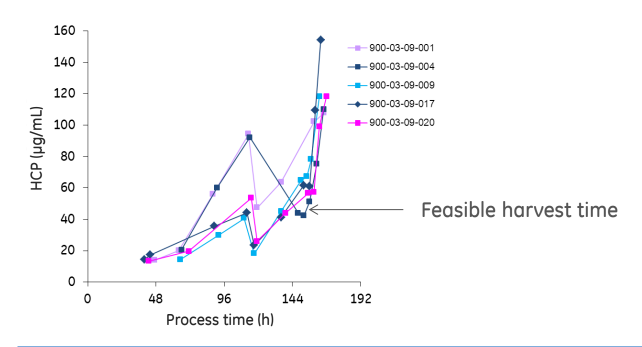


The process consistency was high and virus titers were similar between runs



Virus propagation and release of HCP

HCP content after ELISA analysis



A feasible time for harvest is before the HCP peaks, to facilitate downstream processing





Whole-cell (wP) - Acellular Pertussis (aP)

wP Vaccines

70 year old technology based on killed *B. pertussis* strains

High protection efficiency ~78%

Associated with side effects and safety concerns

The reactogenicity of wP vaccine was thought to be too high to permit routine use in older children, adolescents and adults.

aP Vaccines

Introduced in 1990's

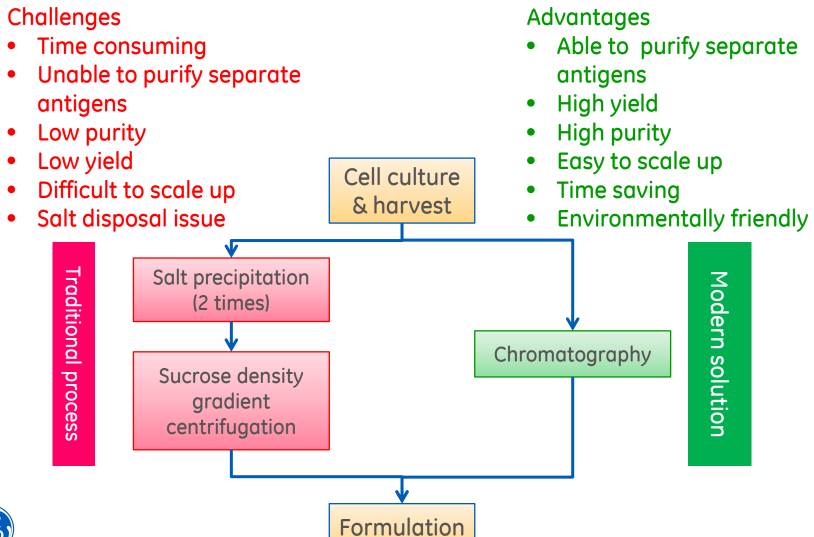
aP contain ≥1 of the separately purified antigens: pertussis toxin (PT), filamentous hemagglutinin (FHA), pertactin (PRN), and fimbriae(FIM) type 2 and 3.

aP is now the dominant type in the industrialized world

aP containing vaccines with reduced concentrations of the antigen have been formulated for use in adolescents and adults



Traditional Process vs. Modern Solution





Project Goal

Traditional process

Chinese pharmacopeia requirement and current situation

- Contain 2 antigens:
 Pertussis toxoid (PT),
 Filamentous Hemagglutinin (FHA)
- Purity >85% (SDS-PAGE)

Yield around 10%

Lack of stable antigen quantitative assay

Current Project

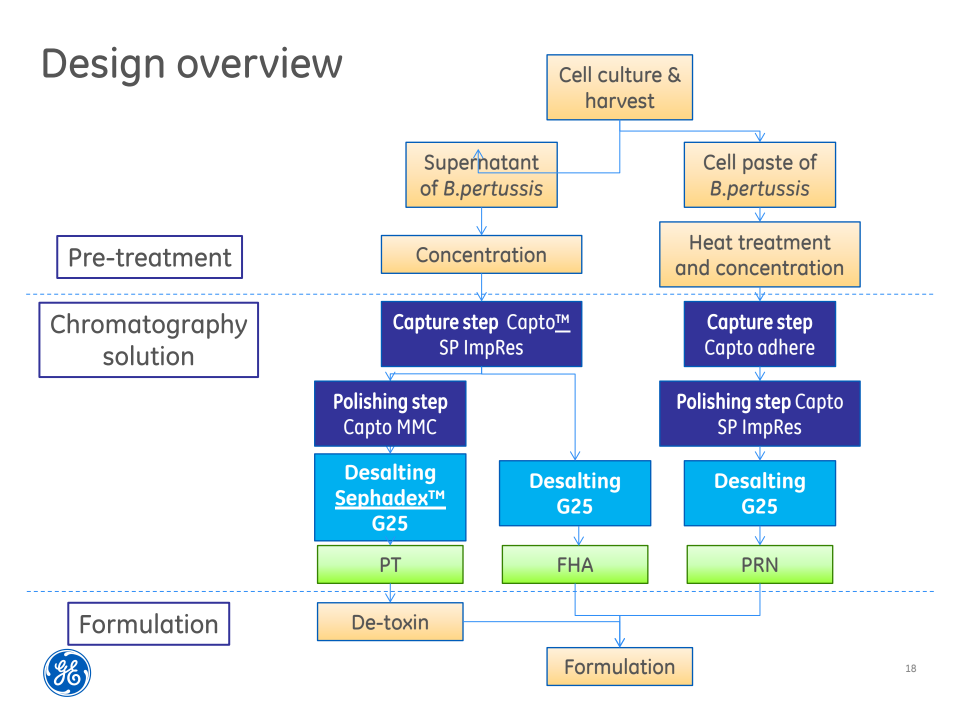
Develop a modern process for pertussis vaccine

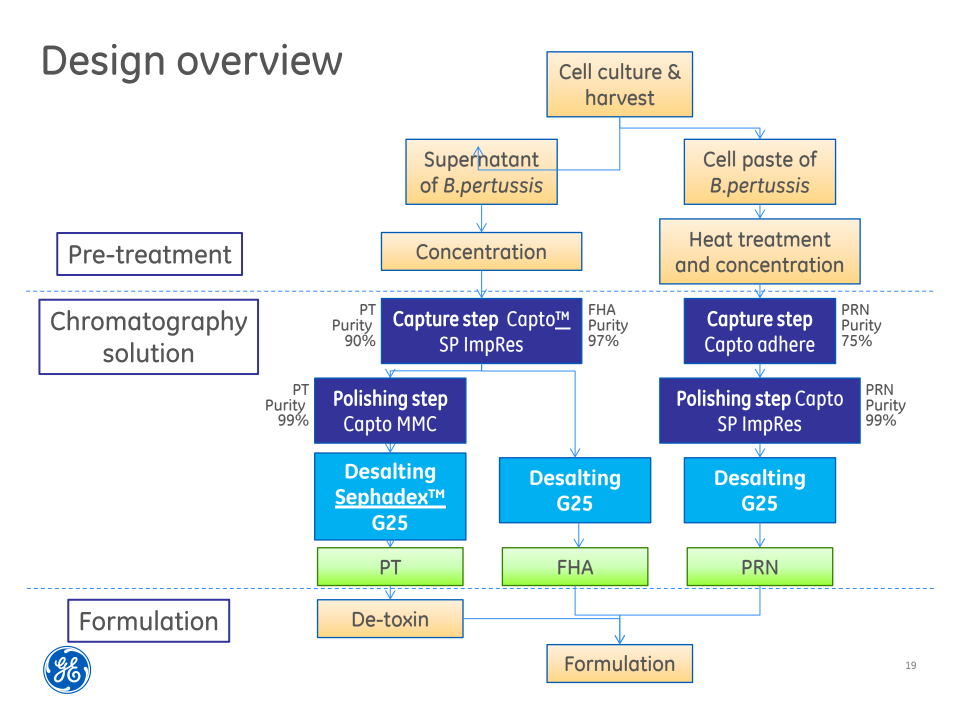
- Contain 3 antigens:
 - Pertussis toxoid (PT), Filamentous Hemagglutinin (FHA) Pertactin (PRN)
- Purity >95% (SDS-PAGE)

Yield >30%

Establish quantitative antigen determination using Biacore™ platform







Process Highlights

- 1. Modern process to produce PT, FHA & PRN using bioprocess friendly, easily scalable, new generation chromatography platform.
- 2. Environmentally friendly.
- 3. Increase purity from 85% to >95%.
- 4. Reduce manufacture time from month to days.
- 5. Recovery increased from 10% to 30%.
- 6. Establish a sensitive, stable platform using Biacore to quantify PT & FHA.





Meningococcal Vaccine

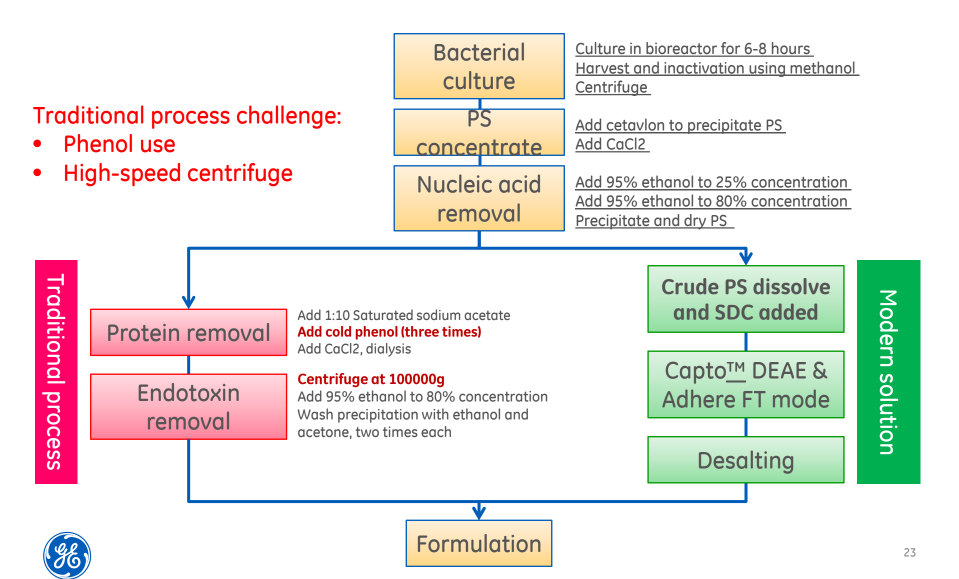


13 clinically significant serotypes. A, B, C, W-135, Y responsible for 90% of global cases

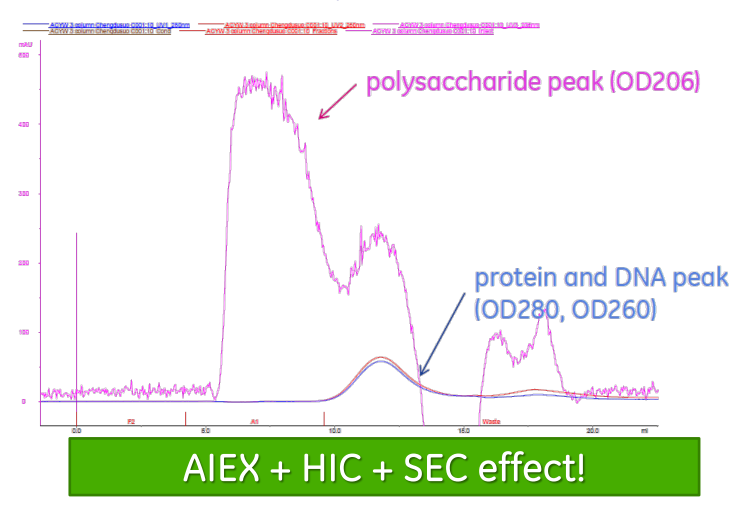
Vaccine for A, C, W, Y are produced using capsular polysaccharide (PS), conjugant technology to enhance immunogenicity



Traditional Process vs. Modern Solution

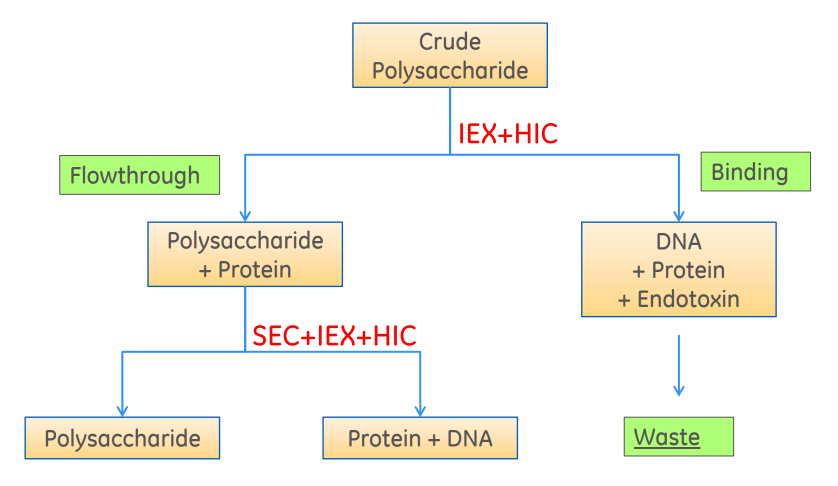


Typical Result From Upgraded Process 1x Capto DEAE + 2x Capto Adhere





Separation Flowchart

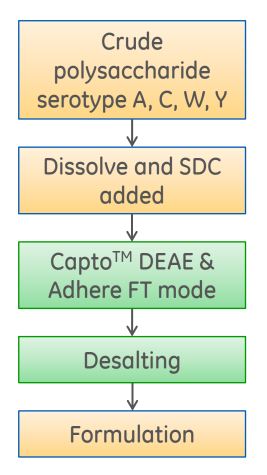




Modern solution for Meningococcal Vaccine A,C,W,Y

Advantages vs. traditional process:

- No phenol use in process, benefit environment & operator's health & safety
- Easy to scale up
- Simple flow-through mode
- All 4 serotypes using same process
- Protein/DNA/endotoxin in products meet requirement

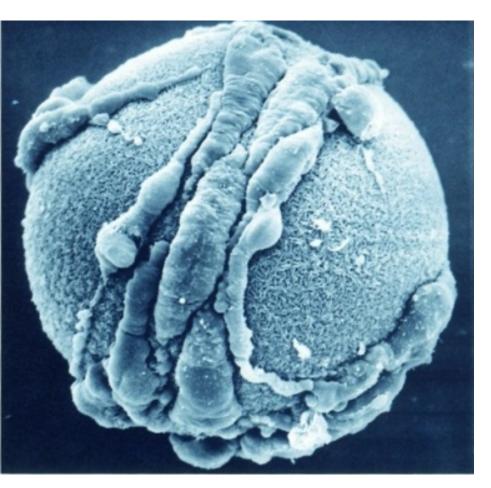




Conclusions



Conclusions



By modernizing legacy vaccine processes there can be improvements in:

Yield

Quality

Scale-up

Cost efficiency

Less hazards



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Thank you!

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