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Design of a multi-stage membrane filtration system for concentration and separation of colloids: example of skim milk microfiltration

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Concentration-separation, example of skim milk microfiltration (0.1 µm)

SP ● ≈ 5 nm

≈ 150 nm

CM

- Microfiltration and ultrafiltration are widely applied in the food sector for concentration or separation of micro or nano-colloids
- Example: MF 0.1 μm of skim milk : Concentration of casein micelles and separation of serum proteins Skim milk = Casein Micelles & Serum Proteins



Industrial design of MF and UF in food sector



Design of multi-stage membrane filtration systems / Objective

> The design rules are not fully established and shared

- The design is made by equipment manufacturers and is mainly based on confidential data and knowhow (equipment manufacturers want to preserve their know-how).

Improvements in the design are still possible

 The design is based on simplifying assumptions (and rarely take into account the relation between the process efficiency and the operating parameters)

> There is a need for optimization of multi-stage membrane filtration design

- To compare various configurations
- To integrate environmental criteria

Propose a general methodology to design the multi-separation-concentration system *Objective 1*: Minimizing the membrane surface for a system operating at desired final retentate (CM) concentration (*VRR*_{out})

Objective 2: Maximising the solute (SP) recovery yield in the permeate



Objective 1 : Required membrane surface area, S



Higher $VRR_{out} \rightarrow \text{lower } J_p \rightarrow \text{higher } S$ S = \$!

S = surface of the shaded rectangle (*Jeantet, Brulé, et Delaplace 2011*)

Objective 1: Interest of a multistage filtration system



 J_{ρ} decreases with $VRR \rightarrow$ multistage system requires less total $S(S_{\Sigma})$ than 1-stage system More stages (N) \rightarrow lower required total S_{Σ} Limitation by the cost of equipment Objective 1: Minimization of total membrane surface area, S_{5}



• Optimal values of VRR_i (i = 1..N-1) for given Q_{in} , $[CM]_{0 \& N}$, and given $J_p = f(VRR)$; • Values of S_i (i = 1..N)



Objective 2: Maximizing the recovery yield of serum proteins in permeate, Y



Recovery yield of SP in permeate Y Y = $[SP]_pQ_p / [SP]_{in}Q_{in}$



Objective 2: Yield of SP vs. configuration of a multi-stage filtration system : Mass balance

$$VRR_{0} = 1 \xrightarrow{VRR_{1}} \xrightarrow{VRR_{2}} \cdots \xrightarrow{VRR_{N}} VRR_{out} = VRR_{N}$$

$$TR_{1}, Y_{1} \qquad TR_{2}, Y_{2} \qquad TR_{N}, Y_{N}$$

Total yield of SP in permeates,
$$Y_{\Sigma}$$

 $Y_{\Sigma} = \sum_{i=1}^{N} Y_{i}$ where $Y_{i} = \frac{1}{VRR_{i-1}} \prod_{j=0}^{i-1} b_{j} - \frac{1}{VRR_{i}} \prod_{j=0}^{i} b_{j}$;
with $b_{j} = \frac{VRR_{j}}{TR_{j}(VRR_{j} - VRR_{j-1}) + VRR_{j-1}}$

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NB: (1)
$$Y_i$$
 depends on all VRR_j and all TR_j ($j = 1..i$)
(2) $TR = f(VRR)$

Objective 2 : Total yield Y_{Σ} vs. TR = f(VRR): Possible scenarios (ex two-stage system)



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What else can affect Jp & TR = f(VRR) and influence the system design ?



3) Concentration of SP in all retentates $[SP]_{r,i}$

- $[SP]_{r,i}$ at VRR_i depends on all VRR_i , j = 1..i
- Influence of $[SP]_r$ on J_p and TR is practically unknown

• $J_{p}(VRR)$ and TR(VRR) at different TMP and v must be measured for given membrane module

Experimental study is needed to elucidate the influence of $[SP]_r$ on J_p and TR

Batch filtration for scale-up of continuous filtration?





Continuous mode vs. batch mode have different $[SP]_r$ at same VRR This can result in different dependencies of J_p and TR on VRR Another reason to verify, whether J_p and TR depend on $[SP]_r$

Example : Microfiltration of skimmed milk; Collection of empirical data

INRAE Dairy platform, STLO, Rennes

- Microfiltration pilot TetraLaval MFS7 equipped with a uniform transmembrane pressure (UTP) system
- **Membrane module** with 7 ceramic membranes Pall 7P1940 GL

membrane length1.02 mnumber of channels19channel diameter4 mmmembrane surface area1.68 m²



- **Milk** (skimmed, pasteurized) provided by Coralis, Cesson Sévigné (F-35)
- Milk enriched with whey proteins concentrate (WPC):

[SP]_{enriched milk} = 2 x [SP]_{milk}

to obtain J_p and TR vs. VRR for two very different [SP]



- *VRR* increased in steps from 1 to 4.2
- Step duration 30 min (tested up to 4h)
- Separate experiments with different combinations of *TMP* = 0.4, 0.7 and 1.0 bar with v = 6.0, 6.5 and 7.0 m/s
- Analysis of [*CN*] and [*SP*] in permeates and retentates

 Determination of
 J_p = f(VRR)
 [SP]_r = f(VRR)
 TR = f(VRR)
 for reference milk and milk enriched
 with SP



Results : $J_p = f(VRR)$ - Permeate flux for reference milk and milk enriched with SP

Low shear and low pressure $v = 6.0 \text{ m} \cdot \text{s}^{-1}$, *TMP* = 0.4 bar For enriched milk + 9.1% of S_{5}

High shear and high pressure $v = 6.5 \text{ m} \cdot \text{s}^{-1}$, *TMP* = 0.7 bar



- Increase of retentate crossflow velocity v and TMP expectedly increase J_p
- $J_{p}(VRR)$ of reference milk > $J_{p}(VRR)$ of milk enriched with SP
- For enriched milk + 9.1% of S_{Σ} is required at low shear/low pressure and + 4.1% of S_{Σ} is required at high shear/high pressure (Hyp: 3 stages, VRR_{out}=3.5, Q_{in} = 1 m³/h)

Results: *TR*=f(*VRR*)- Transmission of serum proteins for reference milk and milk enriched with SP



- At $VRR \leq 2.5$, TR(VRR) of reference milk $\approx TR(VRR)$ of milk enriched with SP
- At VRR > 2.5, TR(VRR) of reference milk strongly decreases
 Increase of [SP]_r with VRR in reference milk did compensate the decrease of TR with increase of VRR
- *TR* expectedly decreased with increase of *TMP* despite of increase of *v*
- For enriched milk + 6.6 % of Y_{Σ} is obtained at low shear/low pressure and + 10% of Y_{Σ} at high shear/high pressure For reference milk, gain in S_{Σ} costs lost in Y_{Σ}

Summary

- A methodology is proposed to design the multi-stage separation-concentration system
 → Theoretical approach (mass balance) + empirical data
- *N*-stage system requires less total membrane surface area S_{Σ} than one-stage system and depends on $J_{p}(VRR)$ Values of VRR_{i} required to minimize S_{Σ} are calculated as $S_{\Sigma,min} = \min_{VRR_{i}} \sum_{i=1}^{N-1} \left(\frac{1}{J_{p,i}} \left(\frac{1}{VRR_{i-1}} - \frac{1}{VRR_{i}} \right) \right)$
- Total yield of permeable component in permeate, Y_{Σ} depends on TR(VRR) and all VRR_i Values of VRR_i required for maximal Y_{Σ} are calculated as

$$W_{\Sigma,max} = \max_{VRR_i} \sum_{i=1}^{N} \left(\frac{1}{VRR_{i-1}} \prod_{j=0}^{i-1} b_j - \frac{1}{VRR_i} \prod_{j=0}^{i} b_j \right) \text{ where } b_j = \frac{VRR_j}{TR_j (VRR_j - VRR_{j-1}) + VRR_{j-1}}$$

- Values of VRR_i required for maximal Y_{Σ} can be different from that required for minimal S_{Σ} (compromise)
- Possible influence of a permeable component concentration in retentate [SP]_r on J_p(VRR) and TR(VRR) must be verified because
 - different combinations of $VRR_{i=1..i}$ can result in different $[SP]_{r,i}$ at same VRR_i
 - batch mode (used for empirical data determination) vs. continuous mode have different [SP]_r at same VRR

Thank you for your attention

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