活性型タンパク質合成のための、再構成型無細胞タンパク質合成系(PURE*frex®*)を用いたアプローチ

Investigation on how to synthesize active proteins by using a reconstituted cell-free protein synthesis system (PURE frex[®])



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<Abstract>

PURE *frex*[®] is a reconstituted cell-free protein synthesis kit developed based on the PURE system. Now the productivity of PURE *frex*[®] is up to 1 mg/mL. However, depending on the protein, it may be difficult to synthesize as functional and soluble form by using only PURE frex[®].

In such cases, we can apply chaperones (DnaK and GroE) for correct folding and isomerases (DsbC and PDI) for disulfide bond formation as additives for PURE *frex*[®]. Here, we introduce examples of protein synthesis that could not be solved by these additives alone, but could be solved by adding other additives or synthesizing at lower temperature.

We mainly show examples of human acetylcholinesterase (hAChE) synthesis. hAChE has three intramolecular and one intermolecular disulfide bonds. When hAChE was synthesized by using PURE *frex*[®] with PDI, active hAChE was obtained but there was a problem in solubility. The synthesis under the presence of surfactant Brij 58 increased the solubility of hAChE but decreased its activity. Next, we examined whether the difference in the solubility of hAChE is depending on the synthesis temperature. The solubility of the product was less than 10% at 30°C although the amount of the product was high. In contrast, the product synthesized at 25°C was almost soluble although the amount was low. It was also found that DnaK/DnaJ/GrpE are essential for the synthesis of soluble hAChE even at lower temperature. This result indicates that it is possible to synthesize difficult-to-express proteins by using PURE frex[®] with optimization of conditions. In this presentation, we also discuss the results from other types of proteins and additives.



2. Result 1: Influence of surfactant on protein synthesis by PURE*frex*®

Experimental conditions					in t	Concentration in the reaction mixture (%)				
for protein synthesis			Surfactant Name	CMC (%)	0.01	0.03	0.1	0.3	1	
	Reaction mixture	Incubation	Template DNA	Digitonin	0.061					
	PURE <i>frex</i> ®2.1 (4 mM GSH) +	37⁰C 4 h	sfGFP PCR product (1 ng/µL)	Sodium cholate	0.628					
				CHAPS	0.492					
				CHAPSO	0.505					
	Sufractants			n-Octyl-β-D-glucoside	0.731					
_	→ Measuremen	nt of GEP fl	uorescence	n-Octyl-β-D-thioglucoside	0.278					
				n-Dodecyl-β-D-maltoside	0.009					
	Most s	urfactar	hte did	n-Decyl-β-D-maltoside	0.087					
	Wost surfactants did			n-Octyl-β-D-maltoside	1.064					
	not inhi	bit the	protein	Mega-8	1.864					
	synthesi	s reac	tion by	Mega-9	0.839					
				Mega-10	0.245					
	PURE <i>fre</i>	x [®] belo	ow the	Triton X-100	0.016					
	CMC			Triton X-114	0.011					
	0110.			NP-40	0.009					
				Tween 20	0.007					
	• Some si	urfactan	ts such	Tween 80	0.002					
				Brij 35	0.011					
	ac Trit/	~ 10	nn and							

 Low temperature
(20°C-25°C)

as	Triton	X-100	and	Brij 58	0.009				
Tween 20 could be used				Fluorescence of synthesized GFP					
even above the CMC.			: CMC (%)	8	80% 50%	10%	1%		
					10	0%: (-) s	urfactant		

3. Result 2: Effect of molecular chaperones on the solubilization of the synthesized product by PURE frex®

3-1. Increasing the amount of DnaK Mix (DnaK/DnaJ/GrpE) facilitates the solubilization of the products during protein synthesis.



- \rightarrow All samples were centrifuged at 20,000 xg for 30 min.
- \rightarrow Synthesized protein and supernatant were applied to SDS-PAGE gel (12.5%).
- \rightarrow Gel staining: Oriole Fluorescent Gel Stain (Bio-Rad)
- \rightarrow Quantitation: LAS-4000 system (GE Healthcare)



3-2. DnaK Mix and ClpB can solubilize the aggregated products after protein synthesis.



4. Result 3: Optimization of reaction conditions for functional protein synthesis by PURE frex®

Human Acet	<u>ylcholinesterase</u>	<u>(hAChE)</u>		
Organism	Homo sapiens			

Basic experimental condition for protein synthesis

Reaction mixture Incubation time **Template DNA**

2

🗕 red

oxi

anti-FLAG WB

Mix (-) DnaK

soluble protein

Mix

PCR product

. (1 ng/μL)

synthesized protein

Synthesized region	32Glu – 614Leu (+FLAG – 6xHis)				
Length	602 a.a.				
Molecular weight	66,812 Da				
No. of disulfide bonds	4 (intermolecular: 1)				

 Measurement of solubilized protein AChE activity assay: Amplite[™] Colorimetric Acetylcholinesterase Assay Kit (AAT Bioquest).

4-1. Influence of surfactants for hAChE synthesis



4-3. It is possible to synthesize soluble and functional

hAChE by synthesis at low temperature.



Most of hAChE synthesized at below 25°C was soluble and had its activity.

4-4. DnaK Mix is necessary for hACHE synthesis even under low temperature condition (25°C).



4-5. Purification of synthesized hAChE

Protein synthesis

Reaction mixture	DnaK Mix	Temperature	Incubation time	Template DNA
PURE <i>frex®2.1</i> (4 mM GSH) 10 μM hPDI/ 0.5 μM hEro1α	(+)	25°C	24 h	PCR product (1 ng/µL)
↓ Purification by Co ²⁺⁻ resin (1 (Binding buffer: 50 mM Tri 5 mM Imi	FALON Meta is-HCI (pH & idazole/ <mark>0.1</mark>	al Affinity Re 8.0)/ 500 mM <mark>% Tween 20</mark>)	sin (Clonte NaCl/	èch))
↓ Activity assay				redhacht
$\vdash F \searrow \Im F = F$				purit
(kDa)		(kI	Da)	-
250*	-	2	50	
150		1	50 —	
100	-	1	00 00	
75		hAChE	75 👝 🛏	
50	•		50 🗕 🛥	
37			37	
25				

5. Summary

- Surfactants can be used for protein synthesis by PURE frex[®]. However, some proteins require investigation in the timing of the use of surfactants. Increasing the amount of DnaK Mix improve the solubility of synthesized PPK2.
- Soluble hAChE is obtained by protein synthesis at lower temperature. DnaK Mix is essential for the soluble protein synthesis even at low temperature.

Optimal conditions for synthesizing proteins with PURE frex[®] differ for each protein. The great advantage of PURE frex[®] is that it is possible to try various variations of additives and synthesis conditions. For more information, please contact us.

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