

## Video Article

# Cell-free Protein Expression Using the Rapidly Growing Bacterium *Vibrio natriegens*

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## Abstract

The marine bacterium *Vibrio natriegens* has garnered considerable attention as an emerging microbial host for biotechnology due to its fast growth rate. A general protocol is described for the preparation of *V. natriegens* crude cell extracts using common laboratory equipment. This high yielding protocol has been specifically optimized for user accessibility and reduced cost. Cell-free protein synthesis (CFPS) can be carried out in small scale 10  $\mu$ L batch reactions in either a 96- or 384-well format and reproducibly yields concentrations of > 260  $\mu$ g/mL super folder GFP (sfGFP) within 3 h. Overall, crude cell extract preparation and CFPS can be achieved in 1–2 full days by a single user. This protocol can be easily integrated into existing protein synthesis pipelines to facilitate advances in bio-production and synthetic biology applications.

## Video Link

The video component of this article can be found at <https://www.jove.com/video/59495/>

## Introduction

Cell-free protein synthesis is a versatile and cost-effective method for the expression of valuable proteins or peptides<sup>1,2,3,4</sup>. Historically, cell-free protein synthesis has been performed using *Escherichia coli* expression systems; however, there has been a recent surge in using alternative, non-model organisms with novel properties as chassis for cell-free expression<sup>5,6,7,8,9,10,11,12</sup>. Organisms with unique metabolic profiles are prime candidates as alternatives to *E. coli* cell-free systems. For example, the marine bacterium *Vibrio natriegens* is the fastest growing of all known organisms with an observed doubling time of less than 10 min<sup>13</sup>. This has garnered *V. natriegens* considerable attention as an emerging microbial host for research and biotechnology<sup>14,15,16,17,18</sup>. Given that the rapid growth rate of *V. natriegens* has been linked to high rates of protein synthesis and metabolic efficiency<sup>19,20,21</sup>, harnessing its cellular machinery for cell-free synthesis may significantly expand the toolkit for rapid protein production and high-throughput screening.

Recently, a cell-free *V. natriegens* expression system has been demonstrated which is capable of producing super folder GFP (sfGFP) at concentrations of > 260  $\mu$ g/mL in 3 h with a T7 promoter<sup>9</sup>. The overall aim of developing this method was to provide users with a highly accessible, cost-efficient, reproducible, and high-yielding cell-free protein expression system that can be prepared using common lab equipment in a short amount of time. This protocol utilizes 1 L cultures in shake flasks, cell lysis by pulse sonication, and small-scale batch reactions in a 96- or 384-well format to maximize parallelization and screening throughput. A long, sustained protein expression is made possible by supplementation of 3-phosphoglyceric acid (3-PGA) as an energy source<sup>8,22,23</sup>. Upon successfully completing this protocol, a user will have the capability to express a desired protein or set of proteins in a cell-free format using *V. natriegens* crude cell extract.

Starting from a glycerol stock, *V. natriegens* crude cell extracts are prepared from cells harvested at an optical density at 600 nm (OD<sub>600</sub>) of 1.0. A 1 L culture will yield approximately 2–3 mL of extract, which is sufficient for more than 800 cell-free reactions at 25% crude cell extract. Proteins can be expressed using plasmid DNA, linear DNA, or mRNA template; however, linear DNA template degradation by endogenous nucleases remains a major drawback when using wild type *V. natriegens* cell-free system<sup>9</sup>. Starting from *V. natriegens* cultures, protein useable for downstream applications can be achieved by a single user in 1–2 full days.

## Protocol

### 1. Preparation of *V. natriegens* Crude Cell Extracts – Bacterial Culture

1. Prepare *V. natriegens* bacterial growth media LB-V2 as per **Table 1**. Sterilize the growth media by autoclaving. Allow media to reach to room temperature (RT). Store excess media at RT.  
**CAUTION:** Wear proper personal protection equipment (PPE) and consult lab specific instructions when operating an autoclave.
2. Use a glycerol stock of wild type *V. natriegens* to inoculate 3 mL of LB-V2 media. Grow overnight at 30 °C while shaking at 225 rpm.
3. Wash 1 mL of the overnight culture by centrifugation at 10,600 x g on a benchtop centrifuge for 1 min. Aspirate the supernatant without disturbing the resulting pellet and resuspend in 1 mL of fresh LB-V2 media.
4. In an autoclaved 4 L baffled Erlenmeyer flask with sterile cover, add 1 L of fresh LB-V2 growth media. Inoculate using 1 mL of washed overnight culture (1:1,000 dilution ratio). Grow culture at 30 °C while shaking at 225 rpm.  
**NOTE:** *V. natriegens* cultures can be scaled up or down while maintaining the 1:1,000 dilution ratio. For example, add 250 µL of washed overnight culture to 250 mL of fresh LB-V2 growth media in a 2 L baffled Erlenmeyer flask with sterile cover.
5. Monitor the culture's OD<sub>600</sub> using a spectrophotometer. When culture reaches OD<sub>600</sub> = 1.0 ± 0.2, harvest culture via centrifugation at 3,500 x g for 20 min at 4 °C. Place pellet on ice.  
**NOTE:** *V. natriegens* grows rapidly, so close monitoring of the culture is necessary. Growth to OD<sub>600</sub> = 1.0 should take approximately 1.5–2 h at the 1:1,000 dilution ratio.
6. Aspirate the supernatant and immediately store the resultant bacterial pellet at -80 °C or directly proceed to cell lysis described in section 2.  
**NOTE:** This step is a good stopping point; however, it is recommended that the pellet be processed immediately or within 1–2 days for best results.

### 2. Preparation of *V. natriegens* Crude Cell Extracts – Cell Lysis

1. Prepare S30 lysis buffer as per **Table 1** in sterile deionized (DI) water and adjust pH to 7.7 using glacial acetic acid.  
**CAUTION:** Glacial acetic acid should be handled with proper PPE when adjusting the pH.
2. Cool S30 lysis buffer to approximately 4 °C in a refrigerator or on ice before beginning the cell lysis procedure.  
**NOTE:** For a quick cool down, lysis buffer can be placed at -20 °C. Do not allow the buffer to freeze.
3. Place cell pellets on ice for 10–20 min or until completely thawed. Resuspend all pellets resulting from the same 1 L culture using 10 mL of cold S30 lysis buffer, then transfer suspension to a 50 mL tube. If pellets are not frozen in the freezer at -80 °C in step 1.6, proceed directly to resuspension.  
**NOTE:** Increase the volume of S30 lysis buffer used to initially resuspend pellets as needed.
4. Centrifuge suspension at 3,500 x g for 10 min at 4 °C. Aspirate the supernatant without disturbing the pellet. Wash the pellet a second time using 10 mL of cold S30 lysis buffer. Place pellet on ice.
5. **In a cold room, add 500 µL of cold S30 lysis buffer to the pellet in the 50 mL tube. Using a wide-bore pipette tip, resuspend the pellet and carefully transfer the entire pellet suspension to a 2 mL tube.**  
**NOTE:** If a wide bore pipette is not available, use a pair of scissors to cut the end of a 1 mL pipette tip to increase the bore for pellet transfer.
  1. Transfer as much pellet as possible without significantly increasing the volume; however, the pellet should be resuspended in enough liquid to be sonicated, as indicated by a homogenous suspension in the 2 mL tube. Do not overfill the 2 mL tube. The suspension should not exceed 1.5 mL; split into multiple tubes if necessary.
6. Keep the cell pellet on ice and work in a cold room. Fill a 600 mL beaker with ice and place a 2 mL tube holder on top of the ice.  
**NOTE:** It is helpful to place the tube holder near the side of the beaker, so the pellet suspension will be visible in the ice to monitor sonication progress (see **Figure 1**).
7. Vortex suspended pellet in 2 mL tube briefly to homogenize the cells, flick tubes to remove any cells on the bottom of the cap, and place into tube holder with cap open. Lower sonicator tip into the suspension so that it is just under the liquid surface.
8. Prepare the sonication set-up as depicted in **Figure 2** using a sonicator and probe with a 1/8-inch tip diameter. Input the following settings into the sonicator control: 20 kHz frequency and 50% amplitude, pulse ON time: 10 s, pulse OFF time: 60 s.
9. Run the pulse sonication protocol for three cycles. If the total volume of the pellet suspension is > 500 µL, run the pulse sonication protocol six times.  
**NOTE:** Generally, 3–6 pulses are sufficient to lyse *V. natriegens* pellets; additional pulses may be required depending on sonicator. During sonication, use the adjustment knob platform to move the probe up and down to lyse any pellet that may have settled to the bottom of the tube. The tube can be removed from the holder and briefly vortexed to re-homogenize the suspension in between pulses. See discussion below for desired consistency of sonicated cells.  
**CAUTION:** Wear appropriate hearing protection when sonicator is active.
10. Following sonication, centrifuge crude cell extract at 16,000 x g for 30–45 min at 4 °C or until lysate is free of any cellular debris as depicted in **Figure 3**.
11. **In a cold room, aliquot 50 µL of the resulting supernatants to new 2 mL tubes without disturbing the pellet.**  
**NOTE:** Any debris that is transferred accidentally into the new 2 mL tubes will drastically reduce extract's capacity for high yielding protein synthesis.
  1. Optionally, set aside 10 µL of cell extract for post-lysis protein quantification in a separate 2 mL tube (see step 2.13 below).
12. Flash freeze crude cell extracts by placing tubes into a tube holder with a dipping string attached as depicted in **Figure 4**. Submerge tubes into a Dewar containing liquid nitrogen and immediately place in a freezer at -80 °C until use.  
**CAUTION:** Use appropriate PPE for handling liquid nitrogen including lab coat, safety glasses, face shield, cryogen apron, and gloves.

- Optionally, quantify the total protein of the cell lysate using 10  $\mu$ L set aside from step 2.11.1 by diluting sample 1:100 in 1x phosphate-buffered saline (PBS) and using any standard total protein quantification assay such as Bradford assay, bicinchoninic acid assay (BCA), etc.

### 3. Preparation of Cell-free Reaction Components

#### 1. General reaction components

- Prepare working stocks of Mg-glutamate and K-glutamate in 50 mL tubes with sterile DI water at concentrations of 100 mM and 2000 mM, respectively.
- Prepare a working stock of 50% (w/v) polyethylene glycol (PEG)-8000 by adding 100 mL of sterile DI water to a 250 mL beaker. Place a small magnetic stirrer into the beaker. Weigh out 50 g of PEG-8000 and add to the 100 mL of water in the beaker.
- Place the beaker on a heated stir plate set to 100 °C and stir at 250 rpm until PEG-8000 is in solution. Allow PEG-8000 liquid mixture to cool before transferring to 50 mL tubes.

#### 2. Energy solution master mix

- Prepare a 5 M solution of KOH by adding 140 g of KOH pellets to 500 mL of sterile DI water.  
**CAUTION:** Prepare this solution in a chemical hood and wear PPE when handling strong bases. The solution will become hot so the bottle cap must be loose to prevent pressure build-up. Allow the KOH solution to reach RT before using.
- Prepare a 1750 mM HEPES-KOH buffer by adding 20.85 g of HEPES to a 100 mL bottle. Slowly add sterile DI water until the volume reaches 40 mL. Vortex bottle to dissolve the HEPES. Use the 5 M KOH solution to adjust the pH to 8.0 and then bring the solution volume to 50 mL.  
**CAUTION:** KOH should be handled with proper PPE when adjusting the pH.
- Prepare the remaining 10x energy master mix stock components at the concentrations indicated in **Table 2** in sterile DI water and place each stock on ice. Thaw the 100 mM ATP, GTP, CTP, and UTP stocks at RT and place on ice.  
**NOTE:** A thermomixer set to 37 °C and 350 rpm can be used to dissolve reagents into solution if necessary. Do not overheat or leave reagents on the thermomixer for an extended period of time.
- In a 15 mL tube, add each energy solution master mix component in accordance to the order and volume specified in **Table 2**. Vortex the solution after each component is added. This will make 5 mL of 10x energy solution master mix.
- Divide the 10x energy solution master mix into 200  $\mu$ L aliquots in 2 mL tubes. Flash freeze each aliquot as performed in step 2.12. Immediately place them into the -80 °C freezer until use.

#### 3. Amino acid master mix

- To prepare fresh 4x amino acid master mix, begin by thawing each amino acid stock at RT and then placing each on ice. Use a vortex and/or thermomixer set at 37 °C and 350 rpm to ensure all amino acid stocks are fully dissolved.  
**NOTE:** Cysteine may not fully dissolve; it can be added to the amino acid master mix as a suspension. Do not overheat or leave reagents on the thermomixer for an extended period of time.
- In a 15 mL tube, add the appropriate volume amino acids to sterile DI water so that final concentration of each is 8 mM in the following order: ALA, ARG, ASN, ASP, GLN, GLU, GLY, HIS, IIE, LYS, MET, PHE, PRO, SER, THR, VAL, TRP, TYR, LEU, and CYS. After adding each amino acid, vortex the master mix solution. The volumes listed in **Table 2** will make up 2.4 mL of amino acid master mix.
- Divide the 4x amino acid master mix into 200  $\mu$ L aliquots in 2 mL tubes. Flash freeze each aliquot as performed in step 2.12. Immediately place into a freezer at -80 °C until use.

#### 4. Production of reaction-ready plasmid DNA template

**NOTE:** Cell-free protein expression in this system has been optimized using the super folder green fluorescent protein (GFP) expression vector T7-pJL1-sfGFP (**Table of Materials**). It is recommended to use this plasmid as a control for cell-free reaction efficiency and the pJL1 backbone for the cloning and expression of other protein sequences. Other plasmid DNA templates can be used; however, it is important to note that transcription is controlled by a T7 promoter sequence and the presence of T7 RNA polymerase. A simple protocol for the large-scale production of any plasmid DNA template from transformed *E. coli* is described below.

- Purify desired vector using a plasmid purification kit as per manufacturer's instruction (**Table of Materials**).  
**NOTE:** Concentrating the plasmid DNA template as much as possible is recommended to meet the tight volume constraints of the cell-free reaction. In general, aim for a 750–1500 ng/ $\mu$ L working stock.

#### 5. Production of reaction-ready mRNA template

**NOTE:** This section is optional. Protein expression has been tested using mRNA template generated from the in vitro transcription of the plasmid T7-pJL1-sfGFP by the listed T7 RNA polymerase (**Table of Materials**).

- Prepare mRNA template from plasmid DNA encoding the protein of interest using the in vitro transcription reaction components in **Table 3**. Incubate reactions for 1 h at 37 °C in a thermocycler.  
**NOTE:** It is recommended to perform 8–10x of these reactions in parallel to generate enough material.
- Pool all transcription reactions. Purify using an mRNA purification and concentrator kit as per manufacturer's instruction (**Table of Materials**). Elute mRNA template into sterile DI water. Store mRNA template at -80 °C until use.

### 4. Performing Cell-free Protein Expression Reactions Using *V. natriegens* Crude Extract

#### 1. Cell-free protein expression using plasmid or linear DNA template

- Remove 10x energy solution master mix and 4x amino acid master mix aliquots from the -80 °C freezer, thaw at RT, and place on ice. Remove the T7 RNA polymerase and RNase inhibitor stocks from the -20 °C freezer and place on ice. Thaw DNA template at RT and place on ice.

2. Prepare a cell-free reaction master mix as per **Table 4** by adding each component in the following order to a 2 mL tube on ice: amino acid master mix, energy solution master mix, Mg-glutamate, K-glutamate, DNA template, PEG-8000, T7 RNA polymerase, and RNase inhibitor. Gently flick the tube after each addition to the master mix.  
**NOTE:** If linear template is to be used for cell-free protein expression, add 5–10x more material as compared to plasmid template to obtain appreciable yields of protein.
  3. Remove *V. natriegens* crude cell lysate prepared in step 2.12 from the -80 °C freezers and place on ice for 10–20 min until thawed. Add the appropriate volume crude cell extract to the cell-free reaction master mix as per **Table 4** and gently mix by flicking or pipetting up and down.
  4. End-point cell-free protein expression using thermocycler
    1. Pipette 10  $\mu$ L of the cell-free reaction master mix to the bottom of a 96- or 384-well PCR plate. In between each transfer to the PCR plate, mix the master mix by flicking the tube gently.  
**NOTE:** Cell-free reaction master mix should be well mixed at all times to maximize reaction reproducibility and cell-free protein expression in all samples.
  5. Briefly centrifuge the plate at 1,000 x *g* for 10 s to pool any master mix that may have become stuck on the sides of the wells. Seal the wells with a plate adhesive to prevent evaporation and then place the PCR plate into a thermocycler set at 26 °C with a heated lid set at 105 °C.  
**NOTE:** The even heat distribution and heated lid of a thermocycler greatly improves protein expression yields.
  6. Incubate the cell-free reactions for a minimum of 3 h. After incubation, expressed proteins can be purified, quantified, and used for downstream processes.  
**NOTE:** Expressed proteins can be directly quantified in the cell-free reaction using a method of the user's choice. For example, fluorescent proteins can be quantified using an external standard curve or radioactivity can be measured if using a radiolabeled amino acid in the cell-free reaction. UV-visible spectroscopy or total protein assays are generally not recommended for directly measuring protein production in cell-free reactions without an initial purification.
  7. Alternatively, monitor cell-free protein expression kinetics using a plate reader.
    1. Pipette 10  $\mu$ L of the cell-free reaction master mix to the bottom of a black 384-well assay plate with clear glass bottoms. Keep the assay plate on ice or work in a cold room while adding the master mix to ensure that the full kinetic profile is obtained. Seal the wells with a clear plate adhesive to prevent evaporation and then place the assay plate into a plate reader set at 26 °C.
  8. Incubate the cell-free reactions for 3–6 h while monitoring the appropriate fluorescent excitation/emission wavelengths corresponding to the expressed protein. For example, monitor at Ex/Em = 485 nm/528 nm for sfGFP.
2. **Cell-free protein expression using mRNA template**
1. Thaw mRNA template prepared in step 3.5.2 at RT and place on ice.
  2. Perform step 4.1.1 through step 4.1.6 as previously specified for linear or plasmid DNA template using **Table 4** to prepare an alternative cell-free reaction master mix for mRNA template.

## 5. Calibration of *V. natriegens* Cell-free Reactions with sfGFP

**NOTE:** This section is optional. The optimal cell-free reaction ion concentration can vary slightly for each crude extract preparation based on the conditions used for cell lysis. Consider performing the optional cell-free reaction ion calibration protocol described below using sfGFP if reaction yields are significantly lower than expected (concentrations < 1.0  $\mu$ g/mL).

1. Prepare the Mg<sup>2+</sup> and K<sup>+</sup> ion calibration solutions as specified in **Table 5** in sterile DI water from the 100 mM Mg-glutamate and 2,000 mM K-glutamate stocks prepared in step 3.1.1. Place calibration solutions on ice until needed.
2. **Following the calibration map in Table 6, pipette 1  $\mu$ L of the Mg-glutamate and 2  $\mu$ L of K-glutamate ion calibration solutions into the appropriate wells in a 384-well PCR plate.**
  1. The order of operations of this step is as follows: Pipette the Mg-glutamate ion calibration solution into the bottom of each well followed by the K-glutamate ion calibration solution onto the side of well without touching the liquid already present in the wells.
  2. Seal the wells by placing an adhesive on top of the plate to prevent evaporation while preparing the calibration cell-free reaction master mix. Gently tap the 384-well plate to mix the Mg- and K-glutamate calibration solutions.  
**NOTE:** A repeater pipette is highly recommended for completing this step reproducibly and quickly.
3. Prepare the calibration cell-free reaction master mix as specified in **Table 5** using T7-pJL1-sfGFP plasmid DNA template in a 2 mL tube. Add each component in the following order to the master mix: amino acid master mix, energy solution master mix, T7-pJL1-sfGFP DNA template, PEG-8000, T7 RNA polymerase, and RNase inhibitor. Gently flick the tube to mix after each component addition.  
**NOTE:** Do not add Mg<sup>2+</sup> or K<sup>+</sup> to the calibration cell-free master mix.
4. Remove the adhesive from the plate. Carefully pipette 7  $\mu$ L of the calibration cell-free reaction master mix on to the sides of the wells without touching the mixed ion calibration solution already present in the wells. Reseal the wells with the adhesive and gently tap the 384-well PCR plate to mix the cell-free master mix with the ion calibration solution.  
**NOTE:** A repeater pipette is highly recommended for completing this step reproducibly and quickly.
5. Briefly centrifuge the plate 1,000 x *g* for 10 s to pool any unmixed liquid that may have become stuck on the sides of the wells. Place the 384-well plate into a thermocycler set at 26 °C with a heated lid set at 105 °C.
6. Incubate the cell-free reactions for 3 h. After incubation, transfer the contents of each well to a black 384-well assay plate with glass bottoms with a multi-channel pipette and read the fluorescence of sfGFP at Ex/Em = 485 nm/528 nm using a plate reader to determine the ion concentration combination that yields the highest amount of protein.

## Representative Results

The described protocol for protein production using *V. natriegens* cell-free expression system can be executed in 1–2 days by a single user, starting from inoculation of culture to protein available for downstream applications. Preparation of crude cell extracts and master mix stocks comprise a significant portion of this time; however, once prepared, most bulk reagents can be stored long term (**Table 7**) and used as needed, shortening the time needed to complete the protocol.

The expression system described is best used with plasmid DNA template or mRNA generated by in vitro transcription reactions. While linear DNA can also be used as template for protein production, it yields significantly lower amount of protein. For example, at optimal reactions conditions at 26 °C, a single 10 µL cell-free reaction can produce > 260 µg/mL of sfGFP in 3 hours using 0.3 pmol of plasmid DNA template or a comparable > 125 µg/mL of sfGFP using 14 pmol of mRNA transcript (**Figure 5A,B**). However, 0.3 pmol of linear DNA will produce significantly less protein (< 20 µg/mL). For each template type, the majority of the protein will be produced within 1–1.5 h; however, it is recommended that reaction is run for a minimum of 3 hours. Cell-free reaction concentrations of sfGFP were determined via linear regression using a purified sfGFP standard curve measuring fluorescence at Ex/Em = 485 nm/528 nm.

The yield of protein production is significantly affected by the concentration of potassium and magnesium ions ( $K^+$  and  $Mg^{2+}$ , respectively). Under optimized conditions, it was found that the optimal  $Mg^{2+}$  and  $K^+$  ion concentrations will be 3.5 mM and 80 mM, respectively (**Figure 6A,B**). Deviations from the optimal ion concentrations may result in a decreased capacity for the cell-free expression system to produce protein with appreciable yields. Additional calibration may be necessary if reaction yields are significantly lower than expected. An optional protocol for ion calibration is described in section 5. This allows for some flexibility in compensating for crude cell extract variability incurred from individual cell lysis equipment and conditions.



**Figure 1:** A close-up view of the beaker and tube holder on the sonication platform. [Please click here to view a larger version of this figure.](#)

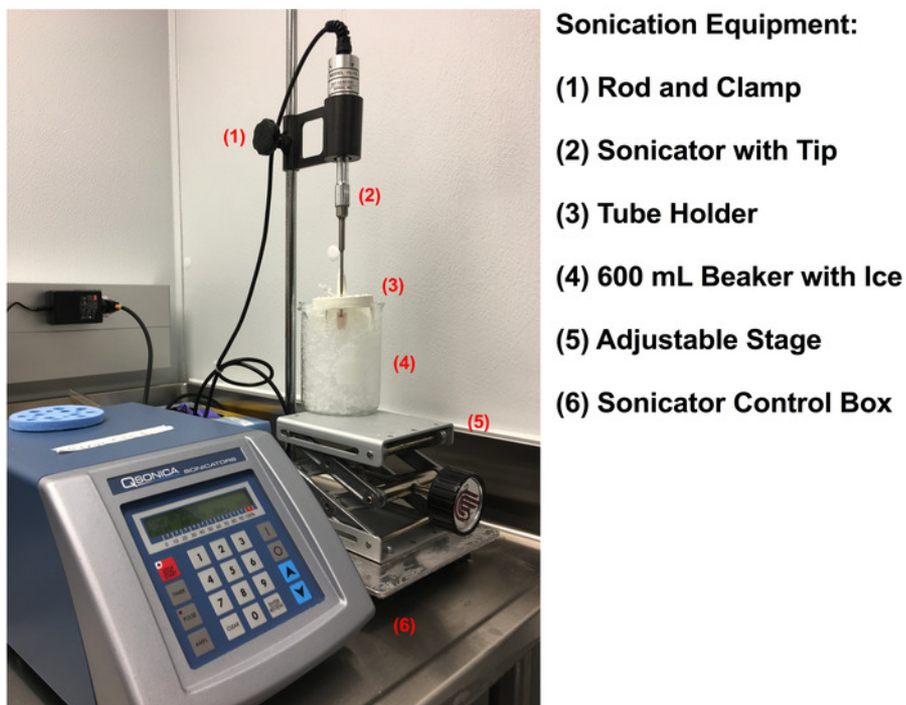


Figure 2: Sonication equipment setup for the preparation of crude cell extract. [Please click here to view a larger version of this figure.](#)

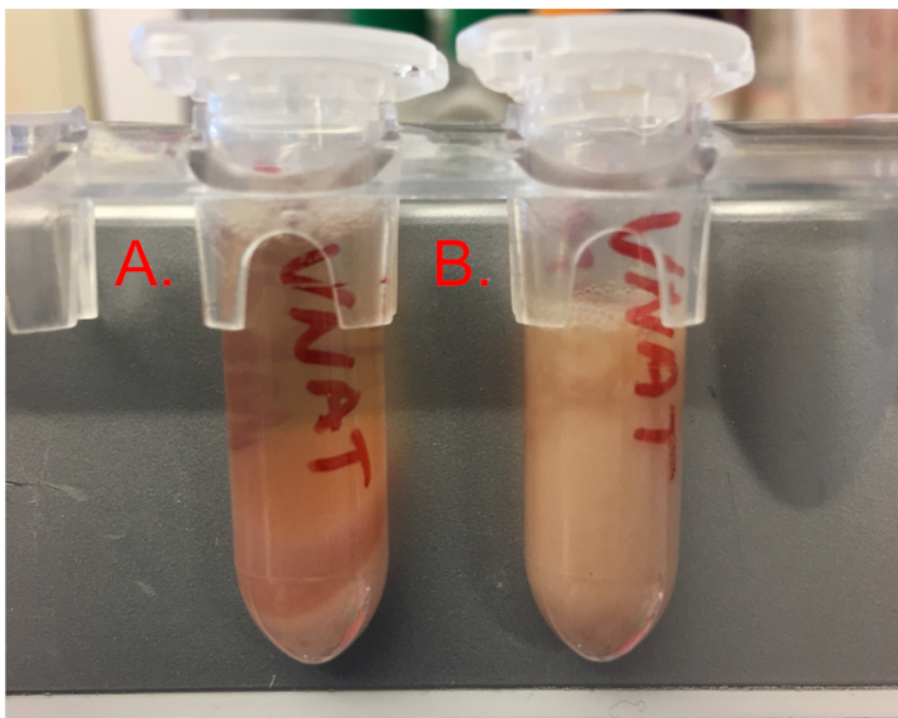


Figure 3: A view of the pellet after sonication and centrifugation. [Please click here to view a larger version of this figure.](#)

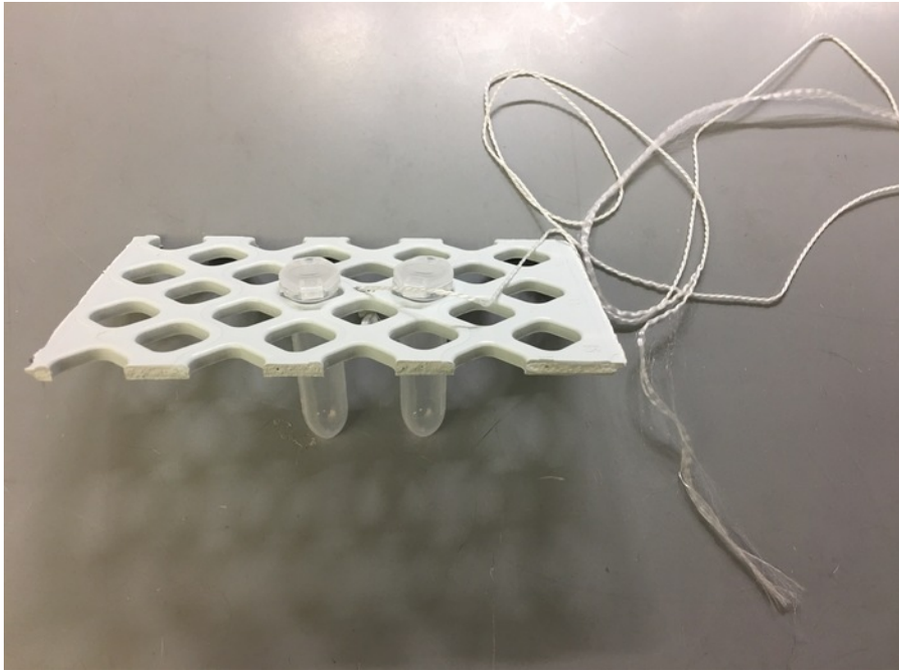


Figure 4: Flash freezing dip for extract storage. [Please click here to view a larger version of this figure.](#)

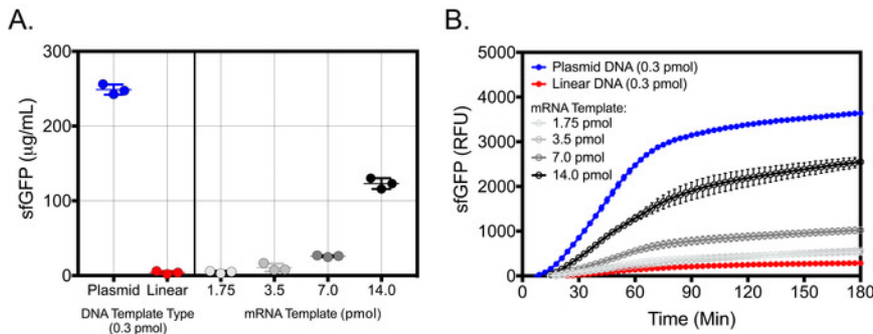
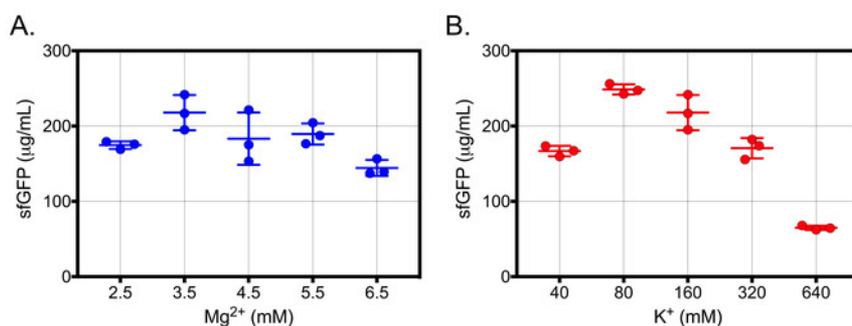


Figure 5: Representative results for *V. natriegens* cell-free protein synthesis using different template types. (A) endpoint assay of sfGFP production using equimolar concentrations of plasmid and linear DNA template (0.3 pmol) as well as increasing concentrations of mRNA template. Cell-free sfGFP concentration was determined using a standard curve of purified sfGFP measured at Ex/Em = 485 nm/528 nm after 180 minutes of incubation at 26 °C at optimal reaction conditions in 10 µL volumes. (B) Kinetic assay of sfGFP production using equimolar concentrations of plasmid and linear DNA template as well as increasing concentrations of mRNA template. sfGFP measurements were taken every 3 minutes at Ex/Em = 485 nm/528 nm over 180 minutes of total incubation time at optimal reaction conditions in 10 µL volumes. For both endpoint and kinetic assays, samples were blank corrected using cell-free reactions supplemented with all components except template. The mean and standard deviations are shown (n = 3). [Please click here to view a larger version of this figure.](#)



**Figure 6: Representative results for ion concentration calibration.** (A) *V. natriegens* cell-free reactions were supplemented with increasing concentrations of Mg<sup>2+</sup> and incubated at 26 °C for 180 minutes in 10 µL reactions. The total concentration of K<sup>+</sup> was 160 mM for all Mg<sup>2+</sup> concentrations. Cell-free sfGFP concentration was determined using a standard curve of purified sfGFP measured at Ex/Em = 485 nm/528 nm. (B) *V. natriegens* cell-free reactions were supplemented with increasing concentrations of K<sup>+</sup> and incubated at 26 °C for 180 min in 10 µL reactions. The total concentration of Mg<sup>2+</sup> was 3.5 mM for all K<sup>+</sup> concentrations. For both calibrations, samples were blank corrected using cell-free reactions supplemented with all components except template. The mean and standard deviations are shown (n = 3). This figure has been modified from Wiegand et al.<sup>9</sup>. Reprinted with permission from Wiegand, D.J., Lee, H.H., Ostrov, N., Church, G.M. Establishing a Cell-Free *Vibrio natriegens* Expression System. *ACS Synthetic Biology*. 7 (10), 2475–2479 (2018). Copyright 2018 American Chemical Society. [Please click here to view a larger version of this figure.](#)

Table 1 A			
Preparation of LB-V2 Bacterial Growth Media			
Component	Quantity (g)	Final Concentration (mM)	Final Volume (L)
LB Broth (Miller)	25		1
NaCl	11.69	200	
MgCl <sub>2</sub>	2.20	23.1	
KCl	0.31	4.2	
Table 1 B			
Preparation of S30A Lysis Buffer			
Component	Quantity (g)	Final Concentration (mM)	Final Volume (L)
Tris Solution (pH 8.0) - 1 M (mL)*	25	50	0.5
Mg-glutamate	2.72	14	
K-glutamate	6.10	60	
Dithiothreitol (DTT) - 1 M (mL)*	1	2	

**Table 1: Reagents for the preparation of 1 L of LB-V2 bacterial growth media and 0.5 L of S30A cell lysis buffer.** [Please click here to download this file in Excel.](#)

Preparation of 10x Energy Solution Master Mix				
Component	Stock Concentration (mM)	Final Concentration (mM)	Quantity (µL)	Final Volume (µL)
HEPES-KOH pH 8	1750	500	1428.57	5000
ATP	100	15	750.00	
GTP	100	15	750.00	
CTP	100	9	450.00	
UTP	100	9	450.00	
tRNA from <i>E. coli</i> MRE 600 (mg/mL)*	100	2	100.00	
Coenzyme A Hydrate	200	2.6	65.00	
NAD	200	3.3	82.50	
cAMP	650	7.5	57.69	
Folinic Acid	100	0.7	35.00	
Spermidine	1600	10	31.25	
3-PGA	2000	300	750.00	
Sterile Deionized Water			49.99	
Preparation of 4x Amino Acid Master Mix				
Component	Stock Concentration (mM)	Final Concentration (mM)	Quantity (µL)	Final Volume (µL)
ALA	168	8	114.3	2400
ARG	168	8	114.3	
ASN	168	8	114.3	
ASP	168	8	114.3	
GLN	168	8	114.3	
GLU	168	8	114.3	
GLY	168	8	114.3	
HIS	168	8	114.3	
IIE	168	8	114.3	
LYS	168	8	114.3	
MET	168	8	114.3	
PHE	168	8	114.3	
PRO	168	8	114.3	
SER	168	8	114.3	
THR	168	8	114.3	
VAL	168	8	114.3	
TRP	168	8	114.3	
TYR	168	8	114.3	
LEU	140	8	137.1	
CYS	168	8	114.3	
Sterile Deionized Water			91.4	

**Table 2: Reagents for the preparation of 5 mL of 10x energy solution master mix and 2.4 mL of 4x amino acid master mix. [Please click here to download this file in Excel.](#)**

T7 RNA Polymerase In Vitro Transcription Reactions				
Component	Stock (mM)	Final Concentration (mM)	Quantity (μL) 1x Reaction	Quantity (μL) 50x Reactions
10x RNAPol Reaction Buffer			1.00	50
ATP	100	0.5	0.10	5
GTP	100	0.5	0.10	5
CTP	100	0.5	0.10	5
UTP	100	0.5	0.10	5
DNA Template (ng/μL)*	1000	500	0.50	25
T7 RNA Polymerase	200	2.6	2.00	100
Rnase Inhibitor, Murine	200	3.3	0.50	25
Sterile Deionized Water			15.60	780
<b>Reaction Volume (μL):</b>	<b>20</b>			

**Table 3: In vitro transcription components for mRNA generation.** [Please click here to download this file in Excel.](#)

Cell-free Reaction Master Mix				
Component	Stock Concentration (mM)	Final Concentration (mM)	Quantity (μL) 1x Reaction	Quantity (μL) 50x Reactions
Extract (%)*		25	2.50	125.00
Mg-glutamate	100	3.5	0.35	17.50
K-glutamate	2000	80	0.40	20.00
4x Amino Acid Master Mix	8.0	2	2.50	125.00
10x Energy Solution Master Mix			1.00	50.00
Plasmid DNA (ng/μL)*	1000	500	0.50	25.00
50% PEG-8000 (%)*	50	2	0.40	20.00
T7 RNA Polymerase			1.00	50.00
RNase Inhibitor, Murine			0.10	5.00
Sterile Deionized Water			1.25	62.50
<b>Reaction Volume (μL):</b>	<b>10</b>			

Alternative Cell-free Reaction Master Mix for mRNA Template				
Component	Stock Concentration (mM)	Final Concentration (mM)	Quantity (μL) 1x Reaction	Quantity (μL) 50x Reactions
Extract (%)*		25	2.50	125.00
Mg-glutamate	100	3.5	0.35	17.50
K-glutamate	2000	80	0.40	20.00
4x Amino Acid Master Mix	8.0	2	2.50	125.00
10x Energy Solution Master Mix			1.00	50.00
mRNA Template (ng/μL)*	2000	4000	2.00	100.00
50% PEG-8000 (%)*	50	2	0.40	20.00
RNase Inhibitor, Murine			0.10	5.00
Sterile Deionized Water			0.75	37.50
<b>Reaction Volume (μL):</b>	<b>10</b>			

**Table 4: Components of optimized *V. natriegens* cell-free reaction master mix for DNA template and mRNA template.** [Please click here to download this file in Excel.](#)

<b>Mg<sup>2+</sup> Calibration</b>		<b>Quantity (μL) 1x Reaction</b>	<b>Quantity (μL) 1x Reaction</b>	<b>Quantity (μL) 100x Reactions</b>	<b>Quantity (μL) 100x Reactions</b>
<b>Final (mM)</b>	<b>Stock</b>	<b>100 mM Mg-Glu</b>	<b>diH<sub>2</sub>O</b>	<b>100 mM Mg-Glu</b>	<b>Deionized H<sub>2</sub>O</b>
2.5	<b>0</b>	0.00	1.00	0	100
3.5	<b>1</b>	0.10	0.90	10	90
4.5	<b>2</b>	0.20	0.80	20	80
5.5	<b>3</b>	0.30	0.70	30	70
<b>Reaction Volume (μL):</b>	<b>10</b>				
<b>K<sup>+</sup> Calibration</b>		<b>Quantity (μL) 1x Reaction</b>	<b>Quantity (μL) 1x Reaction</b>	<b>Quantity (μL) 100x Reactions</b>	<b>Quantity (μL) 100x Reactions</b>
<b>Final (mM)</b>	<b>Stock</b>	<b>2000 mM K-Glu</b>	<b>diH<sub>2</sub>O</b>	<b>2000 mM K-Glu</b>	<b>Deionized H<sub>2</sub>O</b>
40	<b>30</b>	0.15	1.85	12	148
80	<b>70</b>	0.35	1.65	28	132
160	<b>150</b>	0.75	1.25	60	100
320	<b>310</b>	1.55	0.45	124	36
<b>Reaction Volume (μL):</b>	<b>10</b>				
<b>Ion Calibration Cell-free Reaction Master Mix</b>					
<b>Component</b>	<b>Stock Concentration (mM)</b>	<b>Final Concentration (mM)</b>	<b>Quantity (μL) 1x Reaction</b>	<b>Quantity (μL) 50x Reactions</b>	
Extract (%)*		25	2.50	125.00	
Mg-glutamate	Variable	Variable	1.00	50.00	
K-glutamate	Variable	Variable	2.00	100.00	
4x Amino Acid Master Mix	8.0	2	2.50	125.00	
10x Energy Solution Master Mix			1.00	50.00	
Plasmid DNA (ng/μL)*	1000	250	0.25	12.50	
50% PEG-8000 (%)*	50	2	0.40	20.00	
T7 RNA Polymerase			1.00	50.00	
RNase Inhibitor, Murine			0.10	5.00	
Sterile Deionized Water			0.00	0.00	
<b>Reaction Volume (μL):</b>	<b>10</b>				

**Table 5: Ion concentration calibration master mixes.** [Please click here to download this file in Excel.](#)

CF Ion Calibration Map		40 mM K <sup>+</sup>			80 mM K <sup>+</sup>			160 mM K <sup>+</sup>			320 mM K <sup>+</sup>		
		1	2	3	4	5	6	7	8	9	10	11	12
2.5 mM Mg <sup>2+</sup>	A												
3.5 mM Mg <sup>2+</sup>	B												
4.5 mM Mg <sup>2+</sup>	C												
5.5 mM Mg <sup>2+</sup>	D												

Table 6: Ion calibration map. [Please click here to download this file in Excel.](#)

Storage Conditions and Shelf Lives of CF Components		
Component	Storage Location	Shelf Life
Sterile LB-V2 Growth Media	4 °C	3-6 months
<i>V. natriegens</i> Crude Cell Extract	-80 °C	1-3 weeks
100 mM Mg-Glutamate	Room Temp	6 months
2000 mM K-Glutamate	Room Temp	6 months
50% PEG-8000	Room Temp	6 months
10x Energy Solution Master Mix	-80 °C	3-6 months
4x Amino Acid Master Mix	-80 °C	3-6 months
Plasmid/Linear DNA Template	-20 °C	6-12 months
mRNA Template	-80 °C	3-4 weeks

Table 7: Cell-free storage conditions and shelf lives. [Please click here to download this file in Excel.](#)

## Discussion

This protocol has been optimized for wild-type *V. natriegens* and bacterial growth media comprised of LB supplemented with V2 salts (Table of Materials). Other strains of *V. natriegens* can be similarly cultured to generate crude cell extracts for cell-free reactions; however, their use requires additional optimization of this protocol. Additionally, this cell-free protein expression system has been optimized using 3-phosphoglyceric acid (3-PGA) as the primary energy regeneration source. Other energy regeneration sources may be used; however, optimization of reagents and calibration will likely be required to obtain high yielding protein expression<sup>10,11</sup>.

Specific attention to several critical steps in this protocol will ensure maximal extract productivity for high-yielding cell-free protein production. First, crude cell extract must be prepared from *V. natriegens* cell cultures harvested in a mid-exponential phase of growth; protein yield is maximal when cultures reach an OD<sub>600</sub> = 1.0 ± 0.2. While cell-free protein production is possible from cells harvested at a range of optical densities, we have previously found that cells harvested in an exponential state of growth yield significantly more protein<sup>9</sup>. The observed effects of optical density on crude cell extract performance are consistent with those reported for other cell-free expression systems derived from cells grown in batch culture conditions<sup>1</sup>. Because *V. natriegens* grows at a rapid rate, it is critical to closely monitor cultures' optical densities. In general, it is expected that *V. natriegens* cultures should reproducibly reach an OD<sub>600</sub> of 1.0 within 1–1.5 h using this protocol; however, individual growth conditions such as the use of baffled versus non-baffled flasks, which affects aeration, or air versus water incubation, which affects the rate and stability of incubation temperature, may alter growth time. Furthermore, it is generally recommended to culture at least 250 mL of *V. natriegens* in a 1 L baffled flask to ensure a large cell pellet at harvest for easy manipulation and transfer. This greatly improves the success of crude cell extract preparation as well as increases the total volume of extract produced from a pellet. When using smaller scale preparation, culture conditions and reagents can be adjusted appropriately. For large-scale fermentation, further optimization of culture conditions may be required. Finally, to ensure high protein yield, it is critical that cell pellets are processed immediately after harvesting, or within 1–2 days of storing at -80 °C.

The proper lysis of the cell pellet by pulse sonication is critical to the success of cell-free protein expression and is often the most difficult aspect of this protocol for new users. Typically, a well lysed pellet will yield a significant volume of liquid extract that is free of debris. The extract should be slightly viscous but can be easily pipetted when aliquoting into storage tubes before flash freezing in liquid nitrogen. Figure 3 depicts a representation of a well lysed pellet (Figure 3A) in comparison to a poorly lysed pellet (Figure 3B) after the post lysis centrifugation step. A major indication of complete cell lysis is a crude cell extract total protein concentration > 20 mg/mL as determined by a total protein assay (step 2.13). Over-sonication or excessive heating of the crude cell extract will damage the cellular machinery, which cannot be determined without performing a cell-free reaction. Thus, it is highly beneficial to test extract efficiency with a control reaction before dedicating significant time and effort to downstream protein expression applications. While additional optimization may be necessary for different sonication equipment, the pulse sonication steps described have been highly reproducible in our hands.

The use of linear DNA template derived from PCR amplification, restriction enzyme digest, or commercial gene synthesis can significantly increase the capacity for high-throughput and rapid protein production in cell-free expression systems<sup>24</sup>. While protein production from PCR amplified linear template has been demonstrated, the yield of these reactions are approximately 13.5-fold lower compared to reactions using plasmid DNA template at equimolar ratios<sup>9</sup>. This is primarily due to the instability of the linear DNA template which is likely degraded by endogenous nucleases present in *V. natriegens* crude cell extract. While lambda phage protein GamS has been previously used to protect Linear DNA template<sup>24,25</sup>, it was found to be incompatible with *V. natriegens* extracts<sup>9</sup>. Additionally, while increasing the concentration of linear DNA template may allow for a higher protein yield, its fast degradation in crude cell extract will still be a major problem.

A solution to overcoming linear DNA template degradation may be to supplement cell-free reactions with mRNA template generated from in vitro transcription of linear DNA. On the other hand, the use of an RNase inhibitor offers significant protection against mRNA transcript degradation and appreciable protein yields can be obtained in the 10 µL cell-free reaction format (**Figure 5A,B**). Cloning of linear DNA into a circular template through TA ligation, TOPO cloning, Golden Gate assembly, or other recombination methods may be used to circumvent template degradation. Nevertheless, further approaches for inhibition of nuclease activity will be necessary for efficient protein expression using linear DNA template.

To date, several different approaches have been proposed for preparation of crude extract for cell-free protein expression<sup>9,10,11,26</sup>. In developing this protocol, we sought to maximize user accessibility, reduce overall cost, and minimize time-consuming steps. For example, a high protein yield is achieved using a simple two-step sonication-centrifugation process, and does not require cell homogenizers, lengthy dialysis steps or run-off reaction. It is simple to execute in a short period of time and does not require high level of laboratory expertise. Thus, it can help facilitate cell-free expression as a standard for translational academic research and industrial process design.

This protocol expands the toolkit available for investigation and utility of *V. natriegens*, a non-model organism with unique biological properties. Higher protein yield can be achieved by employing semi- or fully-continuous cell-free reactions, to allow for energy regeneration, resupplying of amino acids, and the removal of waste products<sup>3,5,27</sup>. Furthermore, the engineering of wild-type *V. natriegens* to produce DNase- or RNase-deficient strains, removal of deleterious and competing metabolic pathways, and expression of additional tRNAs could greatly enhance the production of proteins in this system<sup>28,29</sup>. As we unravel the biology underlying its rapid growth, further development of *V. natriegens* cell-free systems may accelerate bioproduction capabilities and enable robust expression of therapeutic peptides, small molecules, and synthetic materials.

## Disclosures

DJW, NO, and GMC have filed a patent related to this work.

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