Solid-phase in vitro Transcription and mRNA Purification using Dynabeads[™] Superparamagnetic Beads

Marie Bosnes, Elisabeth Breivold, Laure Jobert, Mu Li¹, Xavier de Mollerat du Jeu¹, Lise Aagaard and Hannah Lindstrøm, ThermoFisher Scientific, Oslo, Norway, 1) ThermoFisher Scientific, Carlsbad, US

ABSTRACT

mRNA-based therapeutics has shown great promise in prevention and treatment of multiple diseases. New tools and scalable workflows for mRNA vaccine development and production will be critical for bringing this promising technology to the clinic. Here we present a proof-of-concept using magnetic beads to simplify the in vitro transcription of mRNA, downstream of mRNA purification.

Streptavidin functionalized Invitrogen Dynabeads™ supermagnetic beads were used for solid-phase in vitro transcription, which generates high quality mRNA, starting with a re-usable biotinylated DNA template immobilized to the beads. The template was easily removed after mRNA transcription by applying a magnet. The immobilized template could be stored for repeated in vitro transcription reactions.

After removal of the magnetic beads, the in vitro transcript (polyadenylated mRNA) was free of template. Further purification and up-concentration was performed by hybridization to Dynabeads™ Oligo(dT)₂₅. providing high quality mRNA with intact poly-A tail. The demonstrated workflow using magnetic beads to immobilize the DNA template, perform in vitro transcription and mRNA isolation, is scalable and easily automated for high throughput and high reproducibility

Bringing mRNA therapy to the clinic will include increasing regulatory requirements. Invitrogen Dynabeads™ have proven track-records with more than 30 years experience in customer partnership and know-how within the field of immune therapy and current Good Manufacturing Practice (GMP).

MATERIALS

- CK19 cDNA cloned into pGEM 4Z-polyA(*) vector
- pCMV-Red Firefly luc plasmid
- pCMV-Red Fireffy luc plasmid DynabeadsTM M-280 Streptavidin (SKU# 11206D, 35136) DynabeadsTM mRNA purification kit (SKU# 65005, 61006) DynabeadsTM MgOne SILANE (SKU# 61005) DynabeadsTM MyOne SILANE (SKU# 37002D) DynabeadsTM MyOne Carboxylic Acid (SKU# 14306D) DynabeadsTM MyOne Carboxylic Acid (SKU# 65012) MEGAscriptTM TT transcription kit (Cat# AM1333) mMESSAGE mMACHINETM T7 ULTRA Transcription Kit (AM1345)

- (AM1345)

- (AM1345) Poly(A) Tailing Kit (AM1350M) MEGAClear[™] Transcription Clean-Up Kit (Cat# AM1908) Invivofectamine mRNA[™] Reagent (custom formulation) BAB/C Mice, female, 4-6 weeks

CONCLUSIONS

- In vitro transcription can be efficiently performed. from a biotinylated template immobilized on Dvnabeads[™] Streptavidin beads.
- Immobilized template can be re-used in sequential in vitro transcription reactions.
- Magnetic removal of template can omit the need for DNase treatment
- In vitro transcribed mRNA purified by Dynabeads[™] Oligo(dT)25, is functional in vivo.
- · Higher concentrations of mRNA can be obtained by including a concentration step, using different types of Dynabeads[™] for generic binding and elution.
- · Prototype beads with higher capacity for mRNA hybridization are under development
- · Magnetic bead handling is flexible, scalable and easy to automate.

ACKNOWLEDGEMENTS

(*) The pGEM4Z-polyA vector was kindly provided by Professor Dr. Joakim Lundeberg, Division of Gene Technology, KTH Royal Institute of Technology, Sweden

RESULTS

Solid-phase in vitro transcription work flow



Figure 1. Solid-phase *in vitro* transcription of template immobilized on Dynabeads M280 streptavidin (a) The vector region spanning the T7 promoter, and the cloned cDNA insert including 30 addinine bases, was amplified by PCR using a biolinylated forward primer and a non-biotinylated reverse primer. (b) The biotinylated template was immobilized on DynabeadsTM A260 Streptavidin and *in vitro* transcription performed by resuspending the beads with template in a transcription mix added T7 RNA polymerase. (c) GeI and (d) histogram showing crude *in vitro* transcriptived (measured by A260 on the NanoDrop One C Instrument) of three 50 µL reactions, using 1 mg of DynabeadsTM A280 Streptavidin immobilized on the VanaDirac based. add 20 µL of the PCR product. Highest yield doisend with lowest truther transcription capoing.



Yield of mRNA in scalable protocols



- ure 2. Purification of mRNA by hybridization to DynabeadsTM Oligo(dT)₂₅ DNA-template was removed from the *in vitro* transcript by applying a magnet, and DynabeadsTM Oligo(dT)₂₅ were added to the *in vitro* transcript in the supernatant. Hybridization was performed for 5 minutes and the beads were washed twice. The mRNA was eluted at 80° C for 2 minutes, and the yield measured by A260 on the NanoDrop One C instrument. Gel showing crude *in vitro* transcript (lane 2), mRNA purified using DynabeadsTM Oligo(dT)₂₅ standard protocol (lane 3), and 10 times increased bead amount in standard binding volume (lane 4). Lanes 1 and 5 are Millenium marker and RibcRuler tadker, respectively.
- (d) Histogram, showing the yield of mRNA isolated using standard protocol and a 10 times direct up-scaling of both bead amount and buffer volumes.

Re-use of solid-phase template in sequential reactions



Figure 3 Re-use of solid-phase template in sequential *in vitro* transcription reactions Set (a) and histogram (b) showing the *in vitro* transcript yield from two templates (A and B) immobilized on 1 m gstreptavidin beads, revised in three sequential 50 µL reactions.

Functionality of oligo(dT)-purified mRNA

(a) In vivo Imaging Systems - 4hr post injection



Up-concentration of captured mRNA using Dynabeads[™] MyOne Silane



Figure 4. Up-concentration of mRNA eluate from Dynabeads[™] Olgo(dT)₂₅ (a) Generic capture workflow, using Dynabeads[™] MyOne SILANE to up-concentrate the mRNA (b) Concentration of mRNA, measured by A260 on the NanoDrop One C instrument, before and after the up-concentration step. The mRNA was from the standard and the up-scaled isolation protocol in Figure 2(d).

(b) Luciferase signal quantification



Figure 5. Functionality comparison between mRNA isolated using different oligo(dT)₂₅ bead prototypes and traditional purification methods In paralell with solid-phase in vitro transcription, a standard in vitro transcription experiment was performed: in vitro valivery and functionality of mRNA purified by prototypes of DynabedsTM M-270 and DynabedsTM M/Q/On Carboxybit, Cold coupled with oligi(dT)₂₅ using proprietary coupling protocols were studied using a luciferase template model. Purified mRNA was intraveneously injected into mice in Invivofectamin, using 0.5 mg/kg and two mice per group.

 demonstrating that all mRNAs had similar activity.
(b) Quantification of the luciferase activity after 4 hrs and 24 hrs, again showing similar activity betwee on nurification methode

CLOSING REMARKS

With more than 30 years experience in customer partnership and know-how within the field of immune therapy and GMP, we enable our customers to develop In the net of minimale therapy and own, we enable our costonie's to be even high quality work-flows. Our tools are based on a comprehensive range of DynabeadsTM monosized superparamagnetic beads, and we offer custom made solutions with beads of different sizes and surface functionalities.



