Testing the Effects of Transfection on Mammalian Cytokine RNA Expression

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Abstract—Abstract outline

This experiment aims to study the effect of genetic expression on polyplex treated cells against β -actin and INF α .

I. INTRODUCTION

Gene therapy is an experimental method for treating disease by introducing a healthy copy of a defective gene into the patient's cells to alter the patients genetic material. The alternative is a non viral gene therapy method which can often have simple and large scale production with low host immunogenicity. However, non viral gene therapy can often yield low transfection efficiencies. This low yield is often due to The inate immune response. The inate immune response to foreign DNA (often by viruses or bacteria) through cytokines as a trigger that rejects the foreign DNA through one of three possibilities. These possibilities include: inducing cell death, halting the production of protien or recruiting immune cells to initiate an adaptive immune response (e.g. antibody protection). New advances in technologies have improved transfection efficiency by modifying cytokines. Cytokines are small protiens that are involved in various types of cell signaling. Classifications of these signalings include interferos, chemokines and interleukins. This experiment aims to see if the inate immune response can be inhibited or manipulated to improve gene therapy treatments.

In this experiment, the prostate cell line will be used for the transfection of foreign DNA. Prostate cells provide a glandular function in the body by generating fluid which serves several functions in reproduction. The analysis will focus on quantifying foreign mRNA that will be produced from the prostate cells. This process will be carried out by nanoparticles that are formed by the self-assembly of DNA/RNA and cationic polymers called polyplexes. Polyplexes are specifically designed to deliver foreign genetic material to cells by a process called transfection.

There are two devices used in this experiment. The first device, the Qubit, is an electronic device that quantifies a substance through fluorescence. The second device is qPCR. Reverse Transcriptase (RT)qPCR is a technique used to describe the level of genetic expression is occuring in vitro by measuring the amount of mRNA within a sample. In this technique, mRNA is reverse transcribed in to complementary DNA by an enzyme labeled reverse transcriptase. For the Real Time PCR experiment, since it is impossible to eliminate all genomic DNA from the cDNA synthesis, two solutions will be prepared. The first solution will be a +RT which

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will include reverse transcriptase and thus, will have cDNA. The second solution will be a -RT solution which will lack reverse transcriptase. When QtPCR is performed on the -RT solution, genomic DNA will be amplified and will be seen in the resulting data. In this experiment, β -actin and RPL13A will be used for direct controls against the immunological transcripts. INF β will not be used as it will only appear after cycle 40 in the experiment producing undesirable results. The No Template Control will consist of DEPC water. The immunological transcripts that will be used in this experiment are Interleukin-6 (IL6) and Inferon- α (INF α) Beta actin is a type of actin isoform which is highly involved in cell motility, structure and integrity. RPL13A is a gene that codes for the 60S ribosomal L13a protein. iCRT will also be present in one polyplex treatment (6hr) to test the effects of polyplex when Wnt pathways (cell surface receptors) are inhibited. Interleukin-6 (IL-6) is a multifunctional cytokine that defends the host in response to immune and hematopoietic activities. Interferon- α is a part of a large subgroup of interferon protiens that help regulate the activity of the immune system.

II. METHODS

A. RNA Purification

The cells in trizol were thawed before phase separation. During phase separation, the poly treated cells in trizol were incubated at 42° after which an 0.2mL of chloroform was added per 1mL of Trizol. The homoginized sample was incubated at 42° for 2-3 minutes following a vigorous shake for 15 seconds. The sample was then centrifuged 12,000 g for 15 minutes at 4°C before transfering the aqueous phase to a second tube. (100Again, the sample was centrifuged 12,000 g for 15 minutes at 4°C . The pellet was not washed in fear of losing RNA. The solution then, underwent a series of centrifuges in between the removal of ethanol to result in an RNA pellet. The pellet was then air dried to remove remaining ethanol. The pellet was incubated at 42° following a resuspension in $20\mu\text{L}$ of RNase free water.

B. cDNA Synthesis

For the first step of cDNA Synthesis, two solutions (+RT/-RT) were formed from the following compounds: 10x dsD-Nase Buffer, dsDNase, Template RNA, polyplex 24hour RNA and nuclease free water. Both solutions were incubated at 42° after being centrifuged. The solutions were then chilled on ice, centrifuged and placed back on ice. For both solutions, 5X Reaction mix and nuclease free water was added. In the +RT solution, Maxima Enzyme mix (reverse transcriptase) was added to the mixture. The -RT solution used DEPC H20

1

instead of Maxima Enzyme mix so that the -RT solution can simulate the +RT solution without synthesizing RNA into cDNA. When the -RT solution undergoes QtPCR, any contaminating genomic DNA will be amplified. The two solutions will be then mixed gently and centrifuged. An incubation period will take place at 25°C for 10 min followed by a second incubation session at 50°C for 15 min. The reaction will be terminated by heating the solutions at 85°C for 5 min. This process was conducted using Maxima First Strand cDNA Synthesis Kit with dsDNase (Thermo Scientific, Cat: K1671)

C. Qt PCR

Each well will contain 10muL of 2X iTag universal SYBR Green Supermix, $5\mu L$ of a primer (IL6, INF α , β -actin or RPL13A), and 5μ L of the respective diluted cDNA. The cDNA needs to be diluted to $4ng/\mu L$ from their respective concentrations before it is added to the wells. The concentrations of cDNA for both sets of Poly24 and no washed cells were all around 100ng/ μ L. Therefore only one set of calculations were needed for determining the proper dilution of cDNA into water for either INF α and IL6 or β -actin and RPL13A and the respective -RT wells. For just cells, a different set of calculations were needed since the original cDNA concentrations were around 11 ng/ μ L. All wells contained the Green Supermix. The columns were sorted in the following manner: 1-3 contained IL6, 4-6 contained INF α , 7-9 contained β -acting and 10-12 contained RPL13A. Refer to Table II for a compressed map of the PCR wells. The QtPCR was ran for 40 cycles.

D. Gel Electrophoresis

In order to further understand the results gained from the QtPCR, an 1.2and gel electrophoresis was ran on the PCR product.

E. Analysis

The results from the QtPCR produced Ct values from each well. A Ct value is a numeric inverse correlation to the quantity of nucleic acid detected by the aparatus. Comparisons are made between Polyplex treated cells for 24 hours version [AV], Polyplex treated cells for 24 hours version [Ts], Polyplex treated cells for 6 hours with iCRT, Cells that were not washed after 6 hours, cells that were not washed and just cells. Each comparison was conducted by taking the difference between the Ct values of the two primers (immunological transcript minus control) on two different treatments. A second difference was taken between these two treatments (i.e. Poly24-NoWash) and then applied to the $\Delta\Delta$ ct method to determine the fold induction. This was repeated three times, and thus an average was determined.

III. RESULTS

A. Qubit

Following the cDNA synthesis, the concentrations were measured by the Qubit. The concentrations are as follows: SD1 (standard 1): $54.27 \text{ng}/\mu\text{L}$, SD2 (standard 2): $1057.52 \text{ng}/\mu\text{L}$, Tk: $8.75 \text{ng}/\mu\text{L}$, DA: $68.0 \text{ng}/\mu\text{L}$, AV: $487.0 \text{ng}/\mu\text{L}$, Ts: $28.0 \text{ng}/\mu\text{L}$.

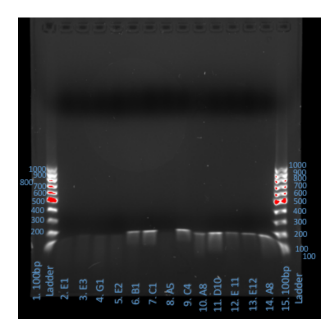


Fig. 1. A 1.2% agarose gel used to further analyze discrepencies seen in the melt curves of the PCR product.

B. QtPCR

TABLE I QTPCR FOLD INDUCTIONS

	IL6- β -act.	IL6-RPL.	INF α - β -act.	INF α -RPL.
Poly24[AV]-NoWash	3.85E-05	7.58E-03	5.09E-05	1.05E-02
Poly24[AV]-Cells	3.10E-02	3.54E-02	3.94E-02	4.27E-02
NoWash-Cells	8.01E+02	4.65E+00	7.01E+02	4.08E+00
Poly24[Ts]-NoWash	8.20E-05	4.64E-03	3.77E-05	6.04E-03
Poly24[Ts]-Cells	6.54E-02	5.90E-02	2.57E-02	2.55E-02
Poly6+iCRT-NoWash6	5.00E+2	4.21E+2	2.58E+1	2.23E+1

Both the Poly24[AV] and Poly24[Tk] variants compared to unwashed cells exhibits fold induction values in the range of 10^{-5} . Processing this value by using a log_2 scale will explain that the polyplex treatment at 24 hours against unwashed cells inhibits genetic transcription levels at a factor of 16.

When comparing Poly24[AV] and Poly24[Tk] variants against washed cells, the resulting fold induction values were at a rate of 10^{-2} . Processing this value by using a log_2 scale will explain that the polyplex treatment in this case against washed cells inhibits transcription levels at a factor of 6.6

The addition of iCRT to Polyplex at a treatment of only 6 hours against unwashed cells at six hours exhibits a fold induction values at a rate of 10^2 . Processing this value using a log_2 scale will explain that the polyplex treatment under these conditions increased transcription levels at a factor of 6.6

As previously explained, when viewing the transcription levels of just cells against unwashed cells, a large difference in transcription levels can be seen.

C. Gel Electrophoresis

Refering to Figure 1, it can be seen that the selections of PCR products was based off of melt curves that were

inconsistent with surrounding wells of similar product (i.e. β actin for just cells). Lanes 2-4 (Wells: E1,E2,E3,G1) consists of -RT solutions. These lanes are empty displaying genomic DNA that is almost undetectable. Lane 8 (Well A5) displays a lack of cDNA for Interferon- α where there should exist amplified cDNA. Lanes 7,9,11 and 14 (Wells: C1,C4,D10,A8) are displaying bands at a greater illumination compared to other lanes. Lanes 7 and 9 consist of Just Cells in Interleukin-6 and Interferon- α respectively. Lanes 11 and 14 are polyplex cells treated for a 24 hour duration in both the [AV] and [Ts] variants for RPL13A and β -actin respectively. Lanes 6,12, and 13 contain smaller amounts of cDNA compared to Lanes 7,9,11 and 14 as they do not contain as great illumination. Lane 6 (Well B1) is cells that were not washed for Interleukin 6. Lanes 12 and 13 (Wells: E11,E12) consist of -RT solutions for RPL13A displaying amplified genomic DNA.

IV. DISCUSSION

The mRNA was purified and converted to cDNA. THe resulting concentration was relatively high compared to peers. This eludes that the treatment for these cells of polycationic DNA for 24 hours could result in higher transcription rates.

Both the Poly24[AV] and Poly24[Tk] variants express higher transcription rates than either just cells or unwashed cells. It can be seen that it depends on both the immunological transcript and if the cell has been washed to what degree.

exhibit small numerical values for fold inductions which displays a large factor of

In the resulting fold induction calculations, it can be seen in Table 1 for both versions of Poly24[AV/Tk] that the treatment of polyplex in cells often yields higher transcription rates when comparing to just cells or unwashed cells.

that in the presence of an immune response the comparison to the

It can also be seen in Table I that the lack of washing cells can diminish the transcription rates.

V. CONCLUSION

To futher understand the nature polyplex treated cells in response to a specific duration, it would be suggested that the polyplex cells are treated in 3 hour increments from 3 hours to 24 hours.

VI. FIGURES

TABLE II PCR WELLS FOR CONTROLS

	Primer	1/4/7/10	2/5/8/11	3/6/9/12
A	Poly24[Tk]	4ng/μL	4ng/μL	4ng/μL
В	Just Cells	4ng/μL	4ng/μL	4ng/μL
C	No Wash	4ng/μL	4ng/μL	4ng/μL
D	Poly24[AV]	4ng/μL	4ng/μL	4ng/μL
Е	-RT	Poly24[Tk]	Cells	Poly24[AV]
Е	NTC	Poly24[Tk]	Cells	Poly24[AV]
G		NoWash(-RT)	NoWash(NTC)	
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TABLE III LIST OF PRIMERS

Primer Name	Tm	Sequence
RTPCR-IL6-FOR	60	CCTTCCAAAGATGGCTGAAA
RTPCR-IL6-REV	60	CACAGCTCTGGCTTGTTCCT
RTPCR-IFNAall-FOR	61	GCACCGAACTCTACCAGCAG
RTPCR-IFNAall-REV	60	ACAACCTCCCAGGCACAA
RTPCR-B-Actin-FOR		TTGCCGACAGGATGCAGAA
RTPCR-B-Actin-REV		GCCGATCCACACGGAGTACTT
RTPCR RPL13A FWD		CCTGGAGGAGAAGAGAGA
RTPCR RPL13A REV		TTGAGGACCTCTGTGTATTTGTCAA

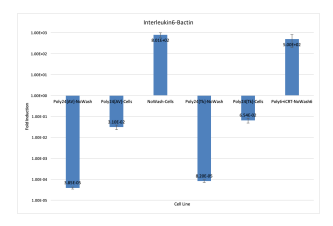


Fig. 2. Logarithmic fold Inductions for Interleukin-6 vs β -actin. Refer to the analysis subsection of the methods section for insight on how this graph was constructed.

REFERENCES

- [1] J. Hagenauer, E. Offer, and L. Papke. Iterative decoding of binary block and convolutional codes. *IEEE Trans. Inform. Theory*, vol. 42, no. 2, pp. 429-445, Mar. 1996.
- [2] T. Mayer, H. Jenkac, and J. Hagenauer. Turbo base-station cooperation for intercell interference cancellation. *IEEE Int. Conf. Commun. (ICC)*, Istanbul, Turkey, pp. 356–361, June 2006.
- [3] J. G. Proakis. *Digital Communications*. McGraw-Hill Book Co., New York, USA, 3rd edition, 1995.
- [4] F. R. Kschischang. Giving a talk: Guidelines for the Preparation and Presentation of Technical Seminars. http://www.comm.toronto.edu/frank/ guide/guide.pdf.
- [5] IEEE Transactions LaTeX and Microsoft Word Style Files. http://www.ieee. org/web/publications/authors/transjnl/index.html

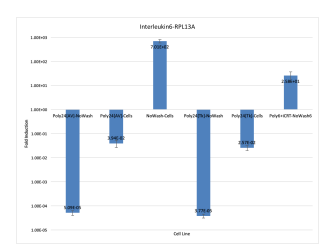


Fig. 3. Logarithmic fold Inductions for Interleukin-6 vs RPL13A. Refer to the analysis subsection of the methods section for insight on how this graph was constructed.

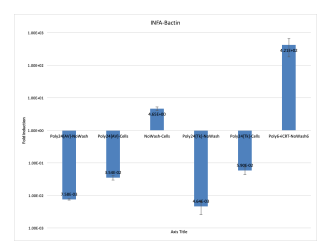


Fig. 4. Logarithmic fold Inductions for Interferon- α vs β -actin. Refer to the analysis subsection of the methods section for insight on how this graph was constructed.

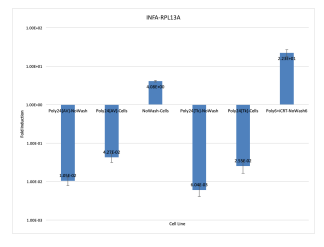


Fig. 5. Logarithmic fold Inductions for Interferon- α an immune cytokine vs RPL13A a standard protien. Refer to the analysis subsection of the methods section for insight on how this graph was constructed.