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The time course of *in vivo* cellular responses to LNPs[†]

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Cells actively respond to drug delivery systems. However, the time course of cellular responses to lipid nanoparticles (LNPs) remains unclear. Here we characterized the transcriptomic response to LNPs carrying mRNA at different timepoints *in vivo*. Exposure to LNPs altered the expression of signaling pathways including endocytosis and lysosomal pathways as soon as one hour after administration. These pathways returned to their baseline state by 24 hours. Our data are consistent with the hypothesis that cells actively yet transiently respond to LNPs.

The FDA has approved three LNP-mRNA vaccines,^{1–3} one LNPsiRNA drug that targets hepatocytes, one GalNAc-ASO drug that targets hepatocytes, and four GalNAc-siRNA drugs that target hepatocytes.⁴ Notably, all six hepatocyte-targeting drugs enter cells through well-characterized mechanisms originally reported in mice,⁵ demonstrating the value of studying the biology of drug delivery. Given the success of these hepatocyte-targeting drugs, there is a growing interest in developing RNA therapies that target new cells and tissues.⁶ These therapies will also likely require scientists to understand the pathways that influence their delivery. One approach scientists have taken is to use RNA sequencing to study the transcriptomic response to LNPs in mice.^{7–9} However, these studies were not designed to reveal the dynamics of that response.

To test the hypothesis that LNPs lead to a dynamic and transient cellular response *in vivo*, we reasoned that it was best to use transcriptomics. Here we used low-input RNA bulk transcriptomic profiling to measure the time-dependent response to nanoparticles, focusing on Kupffer cells. We chose Kupffer cells for two reasons. First, they constitute the largest population of resident tissue macrophages in the body and play a key role in liver function. Second, they readily interact with nanomedicines¹⁰

and affect the efficiency of nanoparticle delivery.¹¹ By measuring the transcriptomic response of Kupffer cells to LNPs one, three, six, and 24 hours after LNPs were administered, we identified time-dependent responses to LNPs. These time-course data also led us to the hypothesis that pre-dosing with the clinically relevant glucocorticoid dexamethasone may improve LNP delivery; subsequent experiments supported this hypothesis.

To assess the temporal response to LNP-mRNA exposure, we employed low-input RNA bulk sequencing technology. We treated mice with either a clinically relevant LNP composed to include the lipid cKK-E12¹² and formulated to carry Cre mRNA,¹³ or a PBS control, and isolated Kupffer cells one, three, six, and 24 hours after injection using fluorescence-activated cell sorting (FACS) (Fig. S1a and b, ESI⁺). We performed low-input SMART-seq v4 using 1000 cells as an input with three replicates per condition (Table S1, ESI[†]). After 72 hours we used flow cytometry to show that the injected LNPs successfully delivered Cre mRNA, with 89% of Kupffer cells (CD45+CD68+) showing tdTomato expression above background (Fig. S1c, ESI⁺). We set a stringent threshold for significance (p < 0.01, fold change >2) for differentially expressed genes; relative to PBS-treated mice, we found 683 upregulated and 2714 downregulated genes one hour following LNP exposure. This total number of genes dropped after three hours, with 1000 up- and 283 downregulated genes after this timepoint, and then 46 up- and 175 downregulated genes after six hours. At the 24 hour timepoint, five genes were upregulated and zero downregulated (Fig. 1a and b). We learned two lessons from these data. First, the number of genes affected by LNPs was high, which is consistent with observations made in previous publications.^{14,15} Second, the maximum number of differentially expressed genes occurred after one hour. These data suggest that cells rapidly respond to LNPs.

We reasoned that RNA sequencing could also quantify the amount of exogenous mRNA we delivered into the cell relative to all the endogenous mRNA made by the cell as it functions. We therefore created a Cre pseudo-gene and quantified its reads at each timepoint. We found that 1.7% of the recorded transcriptome after one hour was exogenously delivered Cre

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Fig. 1 Temporal transcriptomic measurement of the interaction between LNP-mRNA and liver Kupffer cells. (A) Number of genes significantly differentially up- and downregulated compared to the PBS control at each timepoint. (B) Venn diagram comparison of significantly differentially expressed (DE) genes. Normalized exogenous Cre mRNA (C) measured as a percentage of the transcriptome, +/- SD, and (D) ranked in terms of expression.

mRNA (Fig. 1c). This meant that at one hour, Cre mRNA was the sixth most highly expressed gene in the cell (Fig. 1d). This amount of Cre relative to the entire transcriptome decreased to <0.3% at the three- and six-hour timepoints, and at 24 hours the Cre mRNA in LNP-treated mice was similar to the background reads from PBS-treated mice.

After quantifying the number of differentially expressed genes, we hypothesized that they could uncover pathways that were especially affected by LNP delivery. Using the Kyoto encyclopedia of genes and genomes (KEGG) analytical tool, we found that the endocytosis pathway in LNP-treated mice showed a large divergence relative to PBS-treated mice; at one hour there were 53 differentially regulated genes related to endocytosis, and at three hours there were 22 endocytosis-related genes. We noticed similar trends for the phagosome and lysosome; RNA transport, RNA metabolic phases, and RNA polymerase; and the TLR, MAPK, NFκB, PI3K/Akt, and TNF signaling pathways (Fig. S2a, ESI[†]). Of these signaling pathways, TLRs are known to play a crucial role in the innate immune system, and are some of the first immune receptors to interact with LNPs. For example, we previously observed that TLR4 activation dramatically reduces LNP-mRNA delivery.16

The temporal response to LNP exposure can also be seen through gene localization along the TLR signaling pathway (Fig. 2a–c). After one hour, we saw gene colocalization at the surface of the cell (and within endosomes), and genes such as TLR1, TLR6, TLR7, and LBP were significantly differentially expressed (Fig. 2d). Within the cytoplasm, a number of genes linked to signaling pathways considered part of the cascade such as PI3K-Akt, JAK-STAT, and particularly MAPK were stimulated. After three hours none of the surface receptors associated with the TLR cascade were significantly dysregulated, yet 53% of these significantly differentially expressed



Fig. 2 There is a temporal response to LNP exposure within the TLR signaling pathway. (A) Volcano plots of DE genes compared to the PBS control. Genes related to the TLR signaling pathway are in green and quantified in (B). The number of DE genes assigned to the TLR signaling pathway by KEGG analysis peaks after one hour. (C) Heatmap representing the log2 fold change of genes for TLR signaling pathway and where they spatially occur within the cell, relative to PBS one and three hours after injection. (D) Overlaying and colocalizing the DE genes along the TLR signaling pathway after one and three hours, with upregulated genes in green and downregulated in red.

genes are linked to immune response in the nucleus (Fig. S2b, ESI†). Similar temporal cascade effects can be seen in the PI3K-Akt signal transduction pathway, which has been previously linked to nano-particle delivery^{17,18} (Fig. S2c–e, ESI†).

The endocytosis pathway had some of the most striking numbers of dysregulated genes (Fig. S3a and b, ESI[†]), but with a more complex spatio-temporal distribution of where the genes were colocalizing (Fig. S3c and d, ESI[†]). After one hour, we saw gene colocalization around growth factors related to both clathrin- and dynamin-dependent endocytosis (Fig. S3d, ESI[†]). At the endosomal level after one hour, we saw 21 genes downregulated related to early and late endosomes. After three hours, the total number of significantly regulated genes dropped, and where they colocalized on the endosomal pathway shifted.

The genes highlighted in these data should be further studied to understand their impact on drug delivery, with particular emphasis on LNP-specific ligand–receptor interactions that could be involved in LNP uptake and intracellular trafficking (Table S2, ESI†). Analysis of all the differentially expressed genes reveals that only one gene was significantly upregulated at all four timepoints (CCL7). This inflammatory cytokine has also been shown to be upregulated in the thigh muscle fibroblasts of BALB/c mice after intramuscular injection of an LNP with or without an mRNA cargo, indicating a cytokine response at the injection site which depends on one of the LNP lipid components.⁹

The transcriptomic data provided evidence suggesting that pre-treating mice with dexamethasone could improve mRNA delivery to Kupffer cells. Specifically, dexamethasone suppresses TLR1 and TLR6,19 and has also been shown to modulate TLR4 activation,²⁰ which was previously linked to reduced LNP-mRNA delivery.¹⁶ The induction of CCL7 in human peripheral bloodderived mast cells that are stimulated with an anti-IgE Ab has been shown to be significantly inhibited by dexamethasone.²¹ These accumulated data led us to test our hypothesis by measuring mRNA Kupffer cell delivery using flow cytometry at four different LNP doses with a fixed 2.5 mg kg^{-1} dexamethasone pre-treatment one hour prior to LNP administration (Fig. 3a). mRNA delivery increased in Kupffer cells and some other liver cell types (Fig. S4a, ESI⁺). Notably, at 0.3 mg kg⁻¹ LNP-mRNA, we observed a 124% increase in delivery to Kupffer cells in mice pretreated with dexamethasone; we also observed an 82% increase in delivery at the 1.0 mg kg $^{-1}$ dose (Fig. 3a). We then performed a dexamethasone dose response study, and found dose-dependent increases in delivery (Fig. 3b and Fig. S4b, ESI†). Cytokine concentrations in plasma three hours after dosing mice with low and high doses of LNP with and without dexamethasone showed only TNF- α , CCL2 and CXCL10 induced at the 3.0 mg kg⁻¹ LNP dose (Fig. 3c). CCL2 and CXCL10 were inhibited by dexamethasone. These data supported the hypothesis from the transcriptomic profiles that dexamethasone pre-treatment



Fig. 3 Dexamathasone pretreatment improves delivery in Kupffer cells. Pre-treatment of dexamethasone increases aVHH expression in Kupffer cells both at (A) different LNP doses (0.1–3.0 mg kg⁻¹) and (B) different dexamethasone doses (0.1–5.0 mg kg⁻¹). aVHH mRNA was selected to ensure that any observed effects were not driven by the production of a particular protein, thus validating our hypothesis across different functional contexts. (C) Mouse plasma was taken three hours after intravenous dosing. Subjects were pretreated with 2.5 mg kg⁻¹ one hour prior to 0.3 mg kg⁻¹ or 3.0 mg kg⁻¹ LNP. Cytokines were measured by multiplexed ELISA. Two-way ANOVA, **P* < 0.05, ***P* < 0.01, *****P* < 0.0001, average +/– SEM.

could also improve delivery *in vivo*, rather than just improve the anti-inflammatory safety profile. $^{\rm 22}$

In this work, we quantified the transcriptomic response to LNPs at different timepoints as well as the actual amount of exogenous mRNA present in the immune cells. We noted that Cre mRNA was the sixth most highly expressed gene in the cell one hour after LNP administration, constituting 1.7% of the transcriptome. Exposure to LNPs was also shown to significantly alter the expression of the endocytosis and lysosome pathways, as well as important signaling pathways such as TLR, MAPK, and TNF. The temporal differential expression can be seen to progress through the cell with large numbers of genes colocalizing at the membrane level one hour post injection, before the effects being seen in the nucleus three hours post injection. In contrast with other work,¹⁴ no significant pro-inflammatory response is witsnessed at any of the measured timepoints. Examining the transcriptomic profile led us to show that pre-dosing with the clinically relevant glucocorticoid dexamethasone could improve subsequent LNP-mRNA delivery in vivo.

It is important to acknowledge the limitations of this work. First, we did not establish how Kupffer cellular heterogeneity affects delivery; research suggests two distinct Kupffer cell populations.^{23,24} Second, we did not deconvolute the role played by the mRNA cargo and the LNP lipid components by characterizing the transcriptomic response to an LNP without a payload. Therefore, the genes highlighted in this work refer to the *in vivo* cellular response to LNPs encapsulating mRNA. Third, the exogenous mRNA counts represented the biodistribution of the mRNA rather than functional delivery or protein levels. An ideal readout would be able to show the biodistribution of many chemically distinct LNPs, the functional delivery measured as the encoded protein, and the transcriptome of transfected cells.²⁵ Regardless, the list of genes provided in this study may inform future studies as scientists continue to better understand the dynamic responses of cells to LNP-mRNA drugs.

Data availability

The data supporting this article have been included in the main article and as part of the ESI.† All RNA sequencing data are available at SRA (PRJNA1190958). Figure images were created with BioRender.

Conflicts of interest

J. E. D. advises Readout Capital, Nava Therapeutics, and Edge Animal Health. For all other authors, there are no conflicts to declare.

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