Knockdown of *p53* by Accell self-delivering siRNA causes inhibition of p53-dependent DNA damage response in IMR-32 neuroblastoma cell line and β -amyloid toxicity in rat cortical neurons

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Introduction	Experimental design	Testing media compatibility for Accell siRNA delivery in primary neurons
Neuroblastoma cell lines and primary neuronal cultures are commonly used as cellular model systems for studying cancer and neuronal development as well as being highly relevant models for the study of neurodegenerative diseases. However, most neuroblastoma cell lines and practically all primary neuronal cells suffer from low transfection efficiency due to the refractory nature of the cells to lipid-based transfection reagents. As such, application of siRNA for inducing RNA interference (RNAi), has limited utility in these cell types; thus limiting their usefulness for development of functional assays for screening and discovery of novel disease-relevant genes.	Purpose: Determine if knockdown of <i>p53</i> by Accell siRNA will cause suppression of the p53-dependent DNA damage response pathway in IMR-32 cells upon camptothecin treatment. IMR-32 Neuroblastoma cell line Day 0 NTC Accell siRNA	Primary neurons have special media requirements (Neurobasal medium with B27 supplements) so Accell siRNA delivery conditions were optimized to determine the best conditions for cell viability and target gene silencing. Conditions tested: • 100% Neurobasal medium (NBM) • 50:50 NBM and Accell Delivery Media (ADM) • 75:25 NBM and ADM
Dharmacon [™] Accell [™] siRNA enables efficient delivery in a wide range of cell lines and primary cells. Accell siRNA reagents carry a novel chemical modication pattern that facilitates the delivery of siRNA without a peed for	Day 2 Camptothecin	 100% ADM Optimal conditions identied (50: 50 ADM: NBM) that provide requirement eiler size with birth

modication pattern that facilitates the delivery of siRNA without a need for transfection reagents.

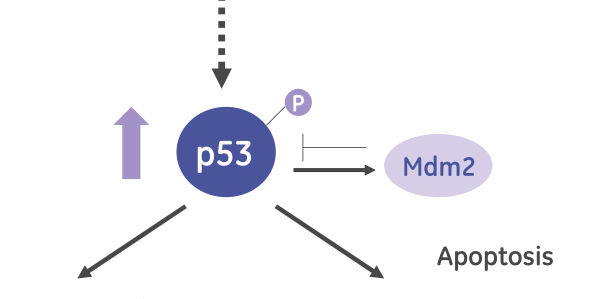
To demonstrate the utility of Accell siRNA reagents in neuronal cells, the effects of the down-regulation of the tumor suppressor p53 was examined. This gene plays a pivotal role in mediating DNA damage-induced apoptosis as well as conferring a protective effect from β -amyloid peptide-induced neurotoxicity. Here we describe how application of Accell siRNA enabled the development of a high content screening assay in IMR-32 neuroblastoma cells and a whole culture cell viability assay in primary rat cortical neurons.

The ability to modulate gene expression in neuronal cell lines and primary neurons using Accell siRNA opens new opportunities for functional genomic siRNA screens in the eld of neuroscience.

p53 is a mediator of numerous cell-damaging agents

Cell exposure to damaging agents induces rapid increase of p53 protein levels in the nucleus, leading to the induction of its transcription targets which control cellular responses such as cell cycle arrest, repair and apoptosis.

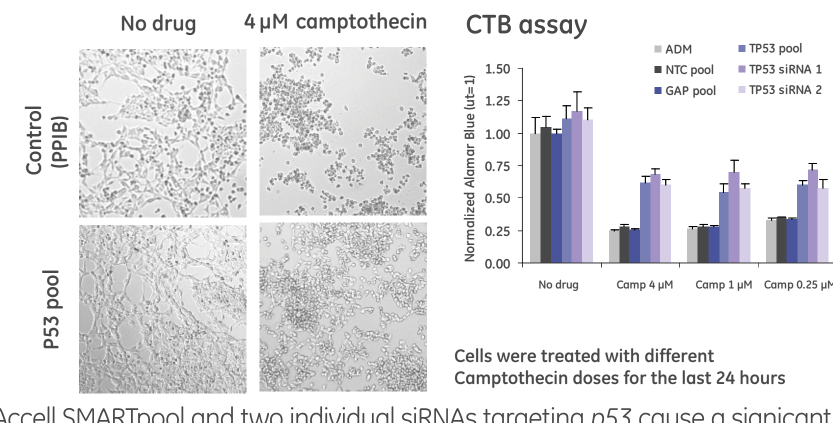
DNA damaging agents/ UV / hypoxia / neurotoxins



Day 3Assay for:
Cell viability
p53 and p21 induction

The cells were analyzed for cell survival by CTB assay and for induction of p53 pathway by High Content Analysis (HCA) using Thermo Scientific[™] Cellomics[™] Multiplexed p53 and p21 Detection Kit.

Knockdown of *p53* increases the survival of IMR-32 after camptothecin treatment



Accell SMARTpool and two individual siRNAs targeting *p53* cause a signicant rescue from camptothecin-induced cell death. This is observed in the phase contrast cell images and the Promega[™] Cell Titer Blue[™] (CTB) cell viability assay at 72 hours post-transfection.

that provide maximum target silencing with high cell viability.



Efficient delivery of Accell siRNA in primary rat cortical neurons

Confocal microscopy reveals virtually all neurons are positive for Accell GAPD Red siRNA (A).
Detailed analysis showed that siRNAs are localised in the

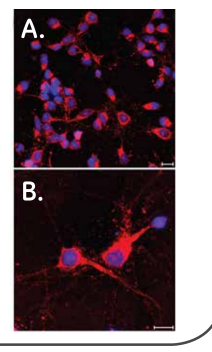
cytoplasm in neuronal cell bodies and in neurites (B).

indicates nuclei. Scale bar 10 µm.

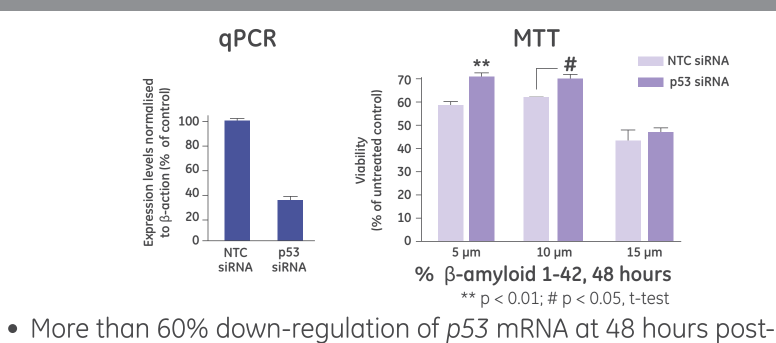
Neurons from E18 rats at 4 DIV (days *in vitro*):

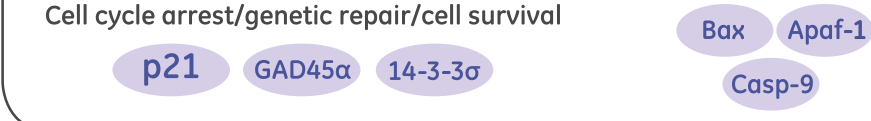
Accell GAPDH Red siRNA (1 µM); 48 hours.

Confocal microscopy (Zeiss LSM 510) Blue staining (Hoechst)



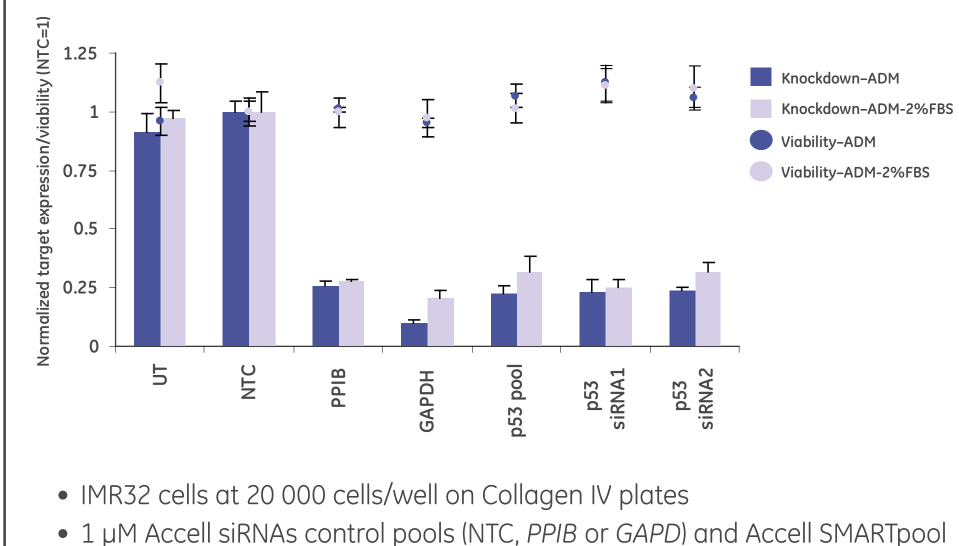
p53 knockdown provides neuroprotective effect from β-amyloid peptide in primary rat cortical neurons





HCA analysis: Reduction in p53 and p21 following camptothecin treatment when *p53* is silenced

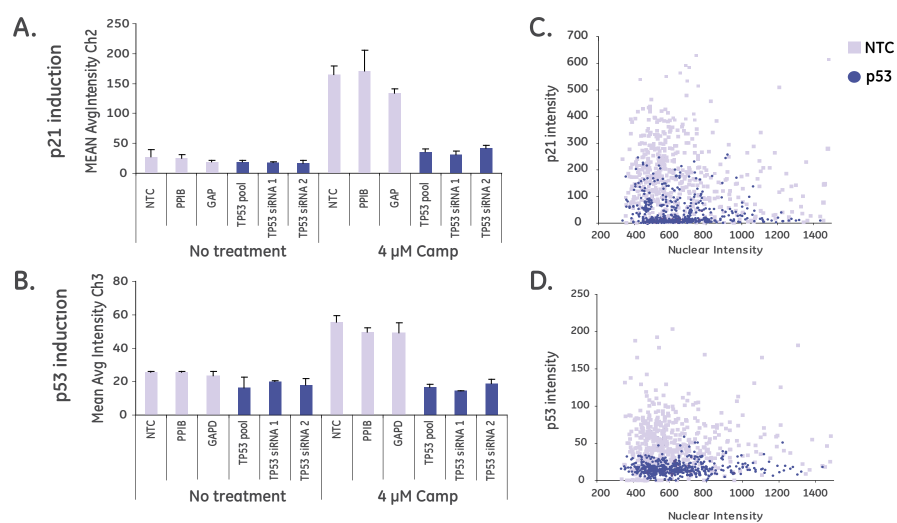
Efficient target mRNA knockdown in IMR-32 cells by Accell siRNA



• 1 µM Accell SIRNAS control pools (NTC, PPIB or GAPD) and Accell SMARTpool or individual siRNAs against *p53* delivered in Accell Delivery Media (ADM) with or without addition of 2% serum

• Knockdown and viability assessed at 72 hours post-transfection

UT = untreated with siRNA; NTC = Non-targeting control pool



Day 0: IMR32 cells transfected with different Accell siRNAs

Day 2: -/+ 4 µM Camptothecin (Camp) for 20 hours

Day 3: cells fixed and stained with Thermo Scientific[™] Cellomics[™] Multiplexed p53 and p21 Detection Kit.

A. Mean nuclear intensities in channel 2 (p21)

B. Mean nuclear intensities in channel 3 (p53)

C. Intensity in channel 1 (Hoechst stain) vs the intensity in channel 2 for p21 nuclear stain (N=500 cell)

D. Intensity in channel 1 (Hoechst stain) vs the intensity in channel 3 for p53 nuclear stain (N=500 cell)

- transfection by Accell SMARTpool reagent.
- Silencing of *p53* resulted in a signicant increase of neuronal survival as compared to NTC siRNA as measured by MTT viability assay.
- The strongest protective effect was observed at 5 μM $\beta\text{-amyloid}.$

Conclusions
 Silencing of p53 in IMR32 cells by Accell siRNA caused an increase in cell survival upon camptothecin treatment.

- HCA with Cellomics Multiplexed p53 and p21 Detection Kit showed a decrease in the p53 and p21 activation following camptothecin treatment in IMR-32 cells transfected with Accell siRNA targeting *p53*.
- Delivery of Accell siRNA was optimized in rat cortical primary neurons.
- Silencing of *p53* in primary rat cortical neurons resulted in a signicant increase of neuronal survival upon β-amyloid treatment.
- Accell siRNA delivery technology permits functional target validation in neuroblastoma cell lines as well as primary cortical neurons.

For more information:

