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Combinatorial CRISPR/Cas9 approaches
targeting different steps in the HIV life cycle
can prevent the selection of resistance

Femke Wolters,
University Medical Center Utrecht,
the Netherlands

nc hiv

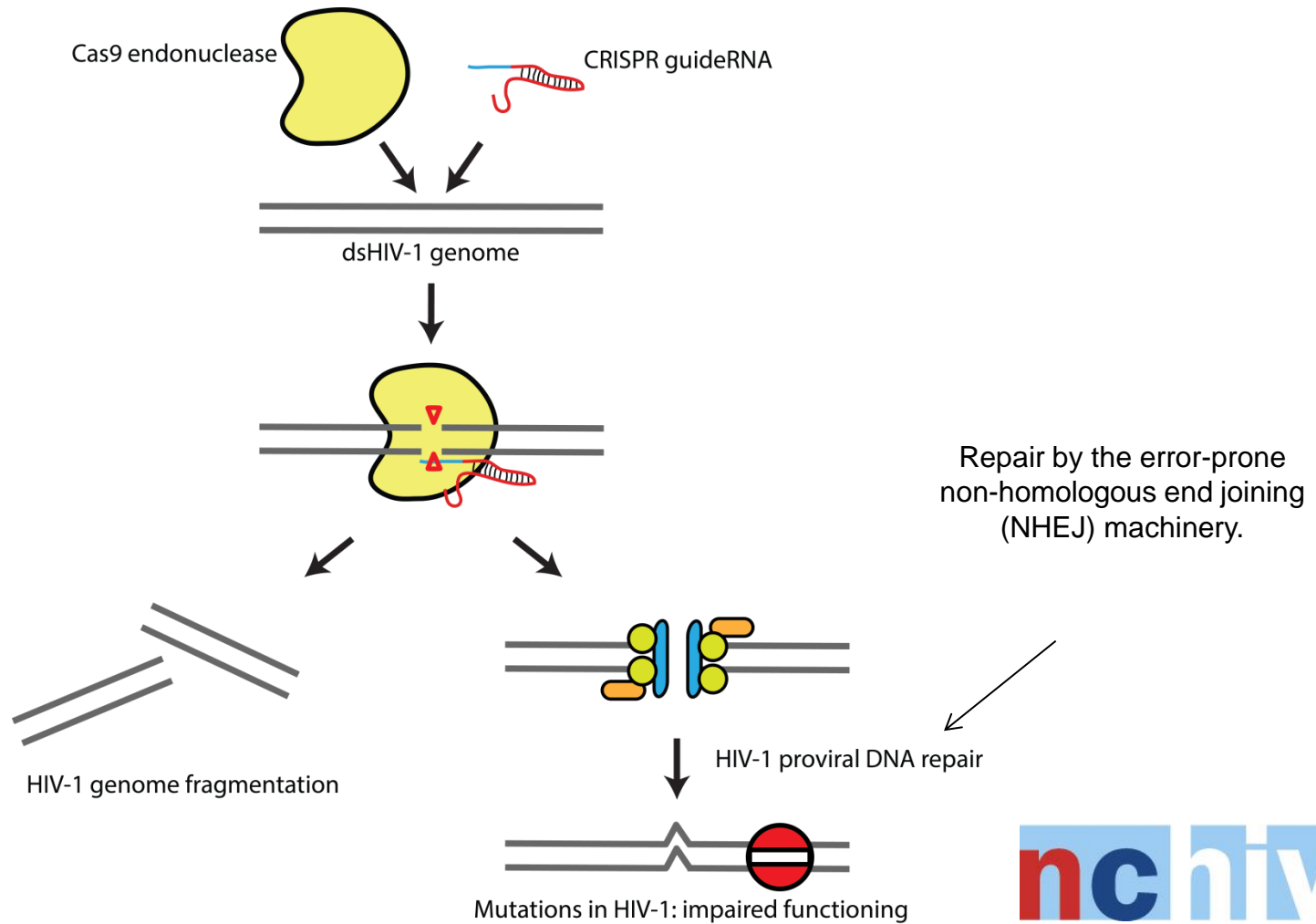
HIV



Current cART strategies

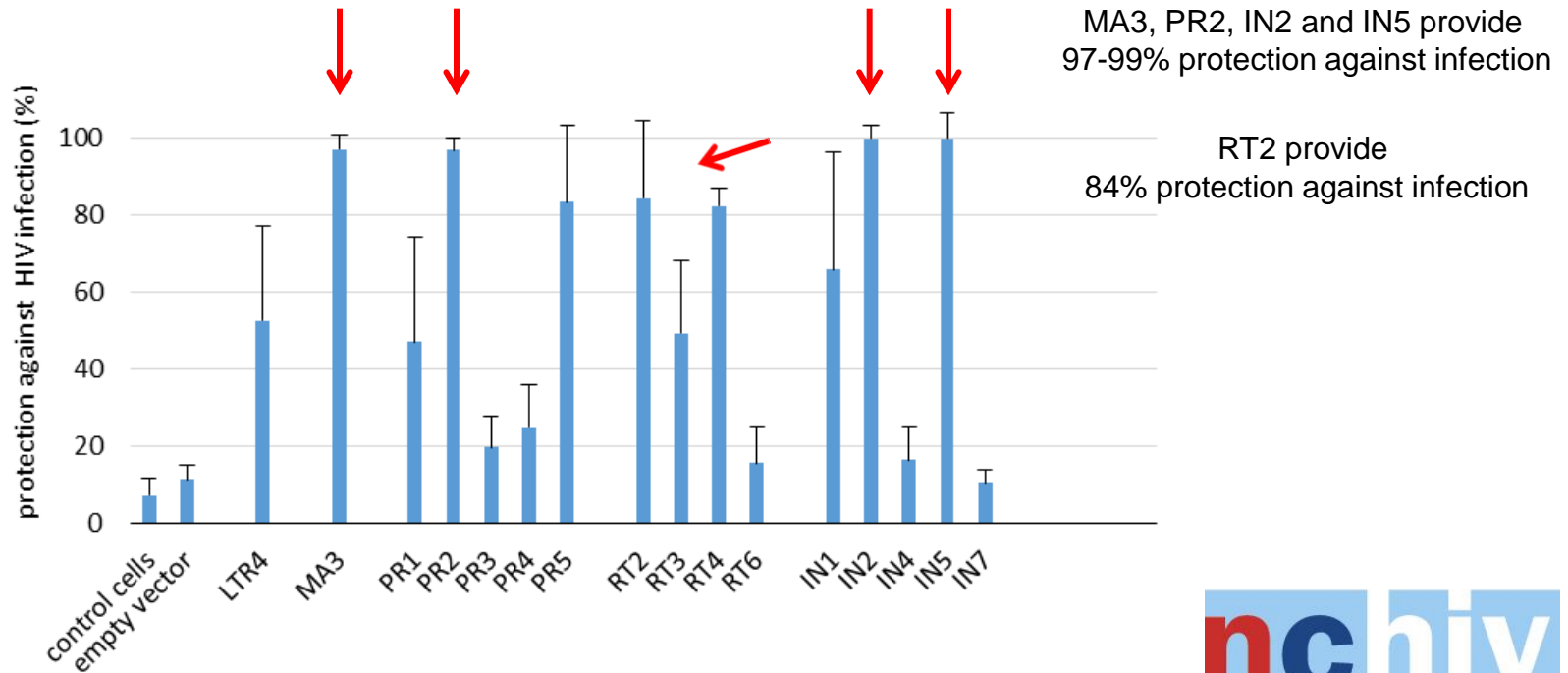
- Current cART is designed to efficiently control HIV replication
- The current arsenal of antiretroviral compounds can neither
- suppress viral production nor target the viral reservoir
- Alternative strategies, such as gene editing, are required to permanently disrupt the HIV genome in the cellular reservoir
- Gene editing tool: RNA-guided CRISPR/Cas9 nuclease

gRNA-guided CRISPR/Cas9 nuclease



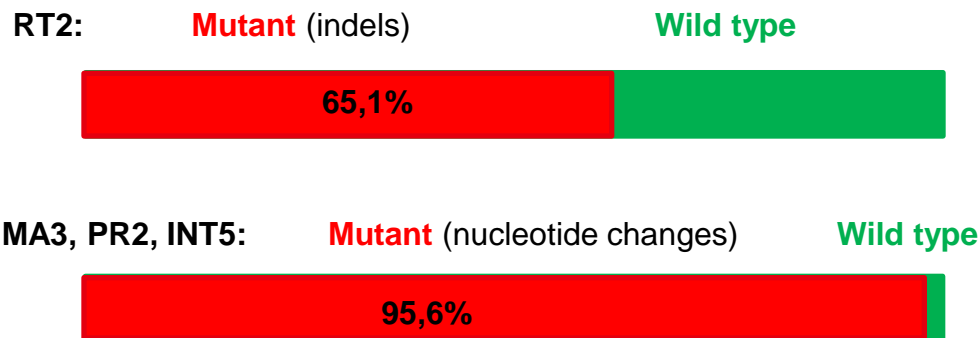
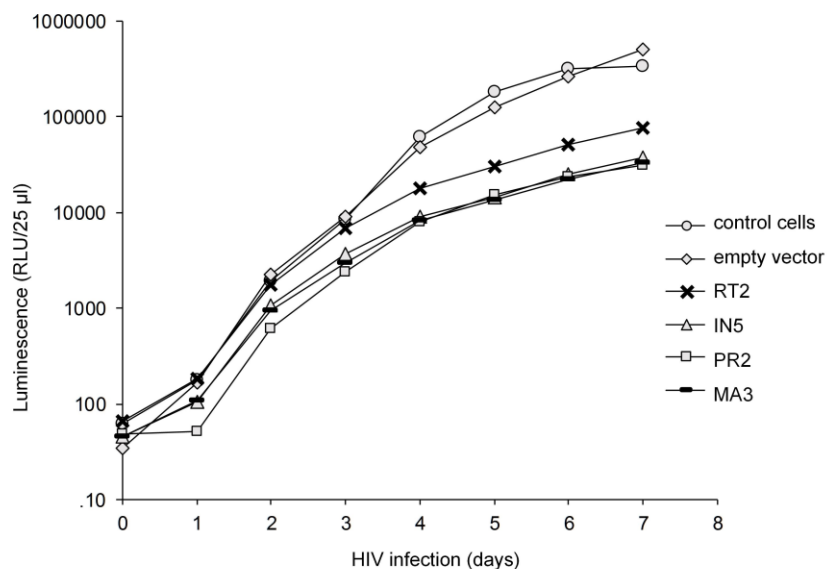
The effect of single gRNAs on viral replication and escape

- Generated T-cells stably expressing single gRNAs and Cas9
- Infect with HIV and measure protection against infection (day 4)



The effect of single gRNAs on viral replication and escape

- Generated T-cells stably expressing single gRNAs and Cas9



- Two recent studies suggest that NHEJ repair machinery may facilitate HIV escape^{1,2}



¹Callaway, Nature News, 2016; ²Liang et al, Retrovirology, 2016.

The effect of single gRNAs on viral replication and escape

Nucleotide substitution patterns in HIV escape variants

HIV target gene Samples (n)	PR 14	RT 21	IN 14	MA 18	Totaal 67
target sequence	ATTAGT	AAGGAA	GTGCTA	GATCGA	
A-->T	4	5	0	1	10
A-->C	2	13	0	0	15
A-->G	10	10	7	10	37
T-->A	0	0*	5	5	10
T-->C	13	0*	5	13	31
T-->G	1	0*	5	18	24
C-->A	0*	0*	1	1	2
C-->T	0*	0*	1	1	2
C-->G	0*	0*	11	0	11
G-->A	2	1	1	1	5
G-->T	1	2	3	1	7
G-->C	1	15	1	1	18
	34	46	40	52	172

Accelerating viral escape

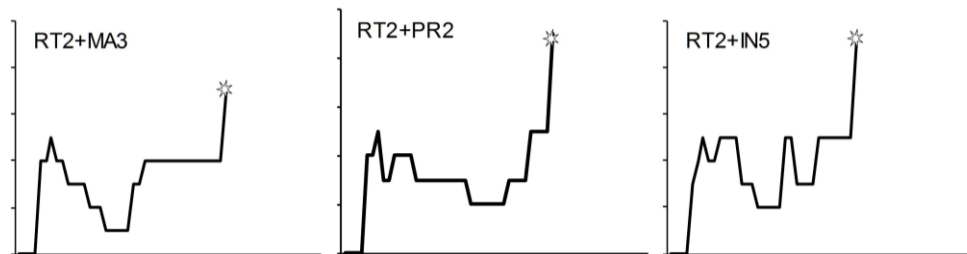
← The cellular error-prone non-homologous end joining (NHEJ) machinery

← HIV-RT and APOBEC3G

* Nucleotide is not present in target site

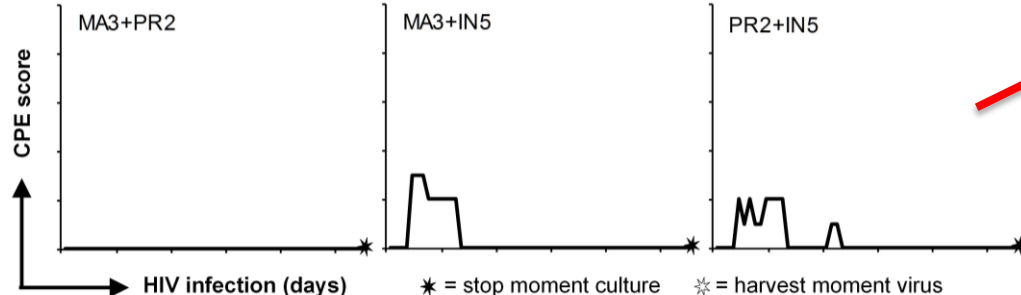
The effect of two gRNAs on viral replication and escape

- Generated T-cells stably expressing two gRNAs and Cas9
- Long-term infection with HIV reporter virus (luciferase) (n=4)



combination of gRNAs	Viral breakthrough (MOI 0.003)				Viral breakthrough (MOI 0.006)			
RT2+MA3	23	34*	38*	-	38*	13	48*	30
RT2+PR2	-	-	-	-	38*	-	-	-
RT2+IN5	-	-	-	23	55*	30*	34*	21*
MA3+PR2	-	-	-	-	-	-	-	-
MA3+IN5	-	-	-	-	-	-	-	-
PR2+IN5	-	-	-	-	-	-	-	-

- = no viral breakthrough
 * = used for deep-sequence analysis of the CRISPR target site





Concluding remarks

- gRNAs differ in potency to inhibit HIV replication
- Rapid viral escape is observed to all single gRNAs
- Error-prone NHEJ repair mechanism is accelerating viral escape
- A combination of two potent gRNAs can successfully prevent viral replication and escape

Gene-editing may provide a future alternative for control of HIV-infection.



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No conflict of interest

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HIV