

Towards a therapeutic HCV vaccine - a preclinical and clinical learning curve

VACCINE TECHNOLOGY II
Albufeira, June 1 - 6, 2008

Alexander von Gabain

Intercell develops *vaccines* 
for the  *prevention and treatment*
of *infectious diseases* .

For more information be invited to: www.intercell.com

Chronic Hepatitis C: Standard of Care

» The treatment of chronic HCV patients is currently based on (pegylated)-**Interferon** and **Ribavirin**

- Significant side effects
- Not all infected patients can be treated
- Significant costs of treatment (up to 30.000 USD per year)
- Long duration (up to 48 weeks)

» Sustained virus response rates are between 50 and 60%, for **genotype 1 only 43-46%** ^{1,2}

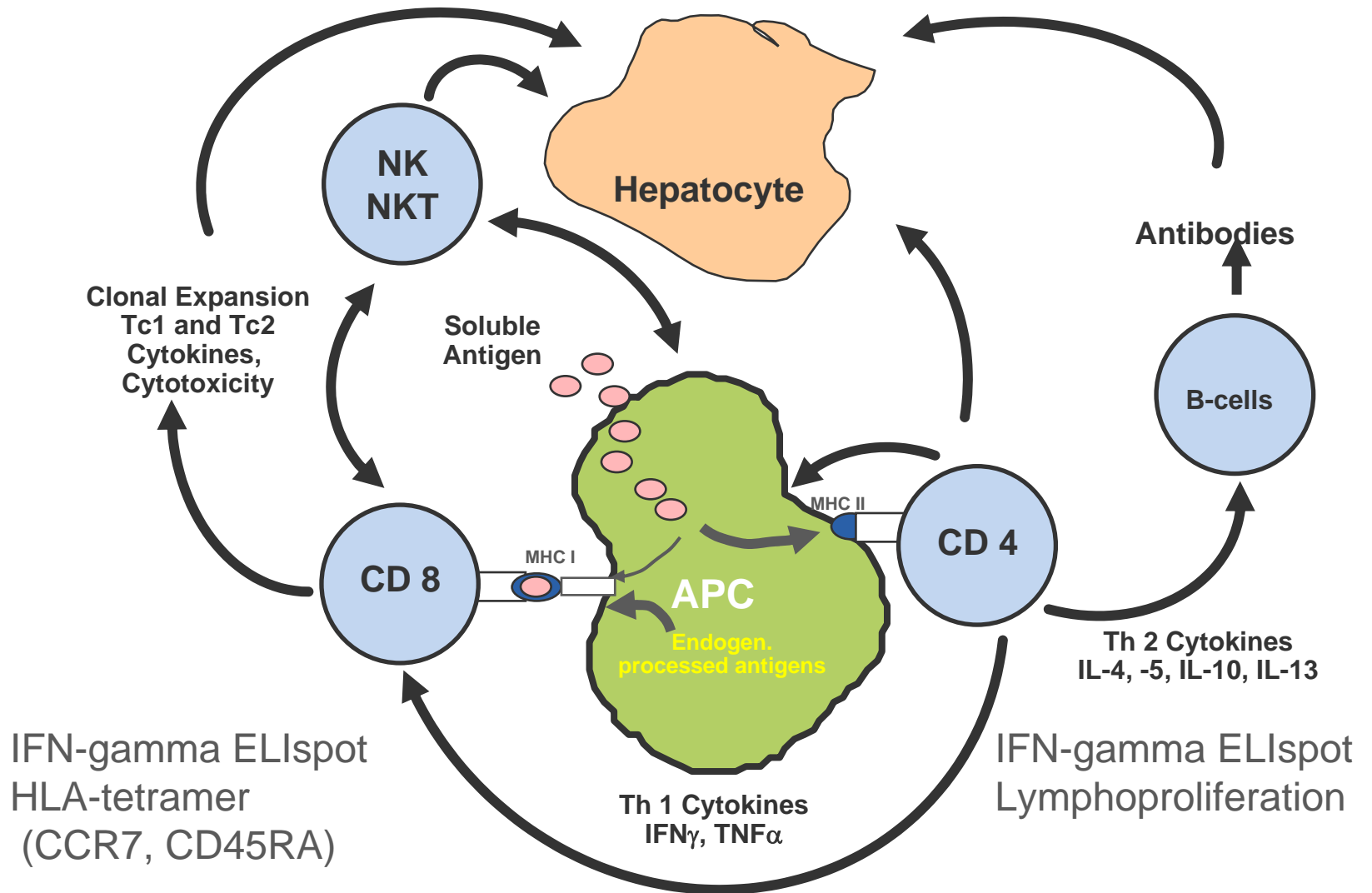
1. Fried M. et al. Peginterferon alfa-2a plus ribavirin for chronic hepatitis C virus infection. N. Engl. J. Med., Vol. 347, 13, 13 Sep 2002.
2. Manns M.P. et al. PIFN alfa-2b plus ribavirin compared with INF alfa-2b plus ribavirin for initial treatment for chronic hepatitis C: a randomized trial. Lancet, Vol. 358 (9286), Sep 2001

HCV: importance of T-cell responses

- » Stronger, broader, quicker and more sustained CD4 and CD8 T-cell responses in self-limited course of acute hepatitis C
- » Response to antiviral therapy may be associated with increased T-cell responses
- » Viral persistence in chronic hepatitis C is associated with immune evasion
 - impaired function of HCV-specific T-cells
 - mutational T-cell epitope escape
- » Chimp models

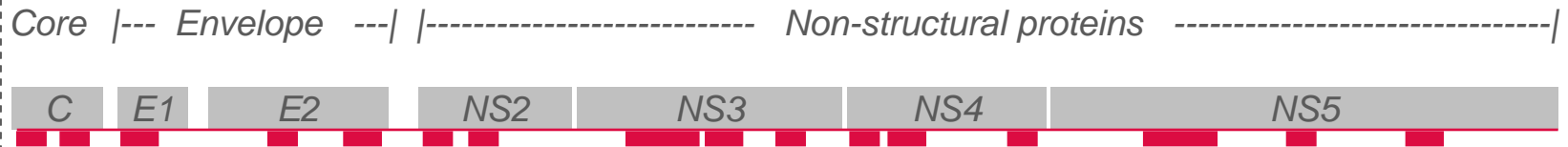
Diepolder 1995, Missale 1996, Rehermann 1996, Lamonaca 1999, Gruener 2000, Thimme 2001, Wedemeyer 2002, Lauer 2002&2004, Cox 2005, Boettler 2005, Spangenberg 2005,...

The T-cell system and Hepatitis C virus infection



The IC41 HCV vaccine: 5 synthetic peptides adjuvanted with Poly-L-Arginine

HCV-
Genome



Intercell
peptide #

Core₂₃₋₄₄

Core₁₃₂₋₁₄₀

NS3₁₀₇₃₋₁₀₈₁

NS3₁₂₄₈₋₁₂₆₁

NS4₁₇₆₄₋₁₇₈₆

Class I
Class II

A*0201

A*0201

A*0201

A*0201

A*0201

DRB1*1101

DRB1*0101

DRB1*0101

DRB1*0401

DRB1*0401

DRB1*0404

DRB1*0404

DRB1*0701

DRB1*0701

DRB1*1101

DRB1*1101

DRB1*1501



 >80% conserved regions in HCV genotypes 1, 2, 3

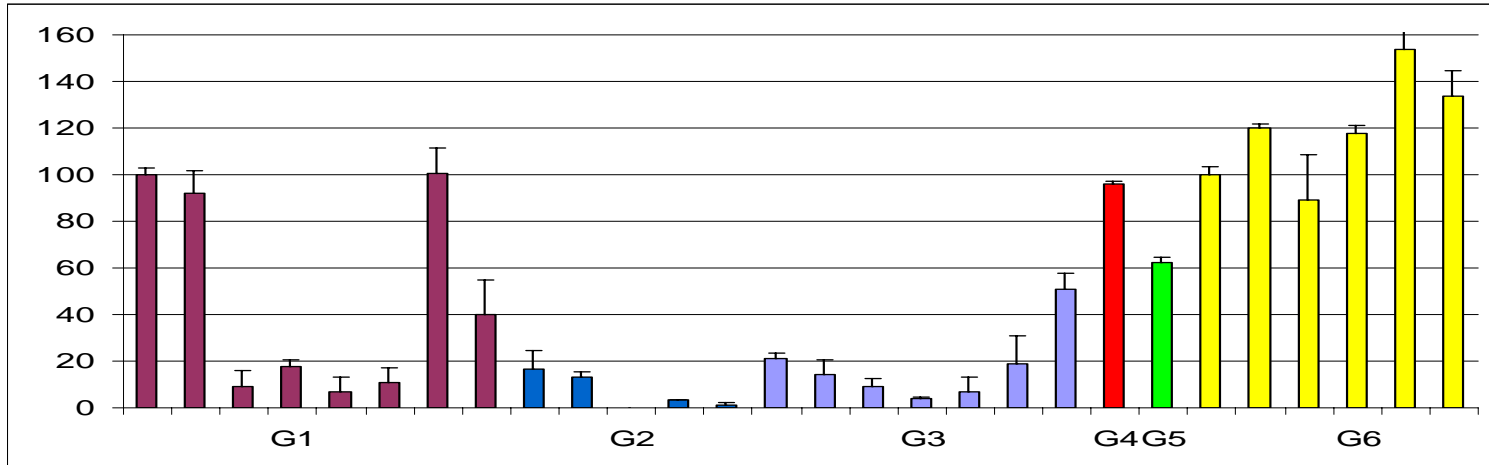
Sequence variability in the NS3-1073 CTL epitope

Position	1	2	3	4	5	6	7	8	9
Wild type	C	I	N	G	V	C	W	T	V
HLA binding		*					*		*
TCR receptor			*		*	(*)	*		
Gen. 1		T	S		A		M	S	I
Gen. 2	T,S		S,A		I	L			
Gen. 3	T,S,A		G	D		T,I			
Gen. 4	A					M			
Gen. 5						M			L
Gen. 6	T,S,A					M,L			

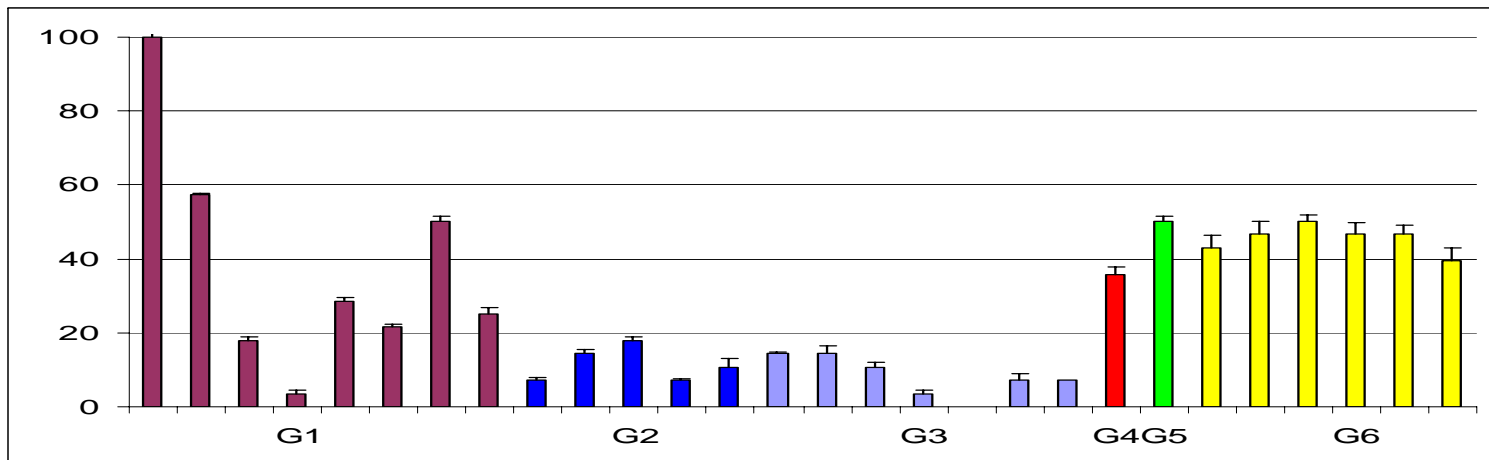
- ➔ **Conservative (green) and non-conservative (red) amino acid exchanges** in each position of the NS2-1073 peptide among the different genotypes of the Hepatitis C Virus.
- ➔ * indicates the positions important for HLA binding or for the TCR receptor recognition.

Cross-genotype recognition of twenty-eight NS3-1073 peptide variants

IFN- γ ELISPOT USING T-CELLS INDUCED AGAINST WILDTYPE



In vitro T-cell line



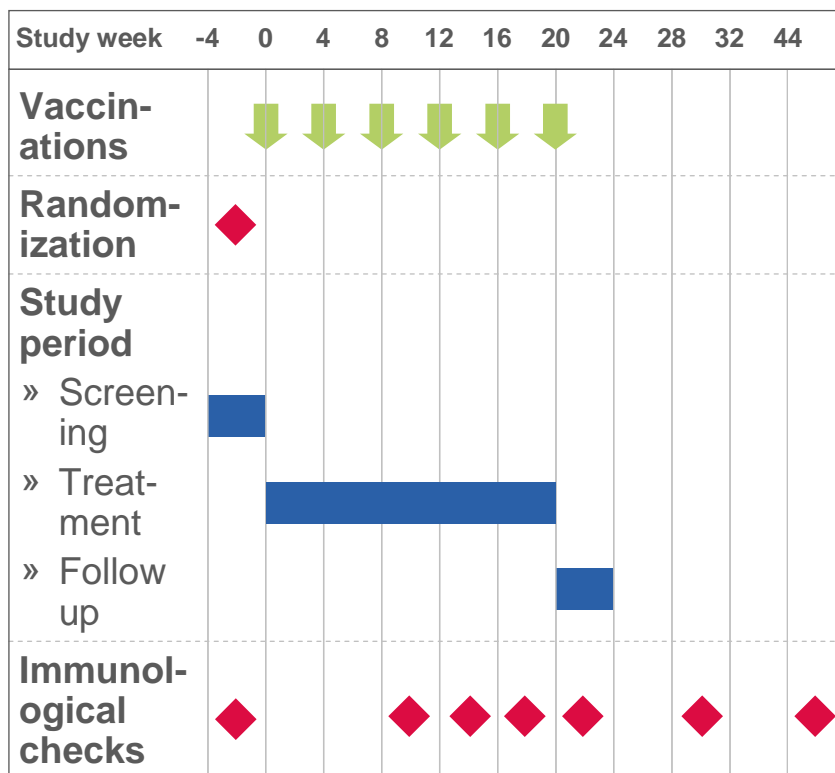
Ex vivo Elispot IC41 vaccinated healthy volunteer

IC41-1: 60 chronic HCV patients, standard IFN/riba therapy non-responders/relapsers

Phase II in non-responders

TREATMENT SCHEDULE AND STUDY DESIGN (IC41-201)

Treatment schedule



Study design

		5 Hepatitis C Peptides**	Poly-Arginine**	No. of patients
Control groups	B	0.00	2.00	12
	C	5.00	0.00	12
Treatment groups	G	2.50	1.25	12
	H	2.50	2.00	12
	K	5.00	2.00	12
Total				60

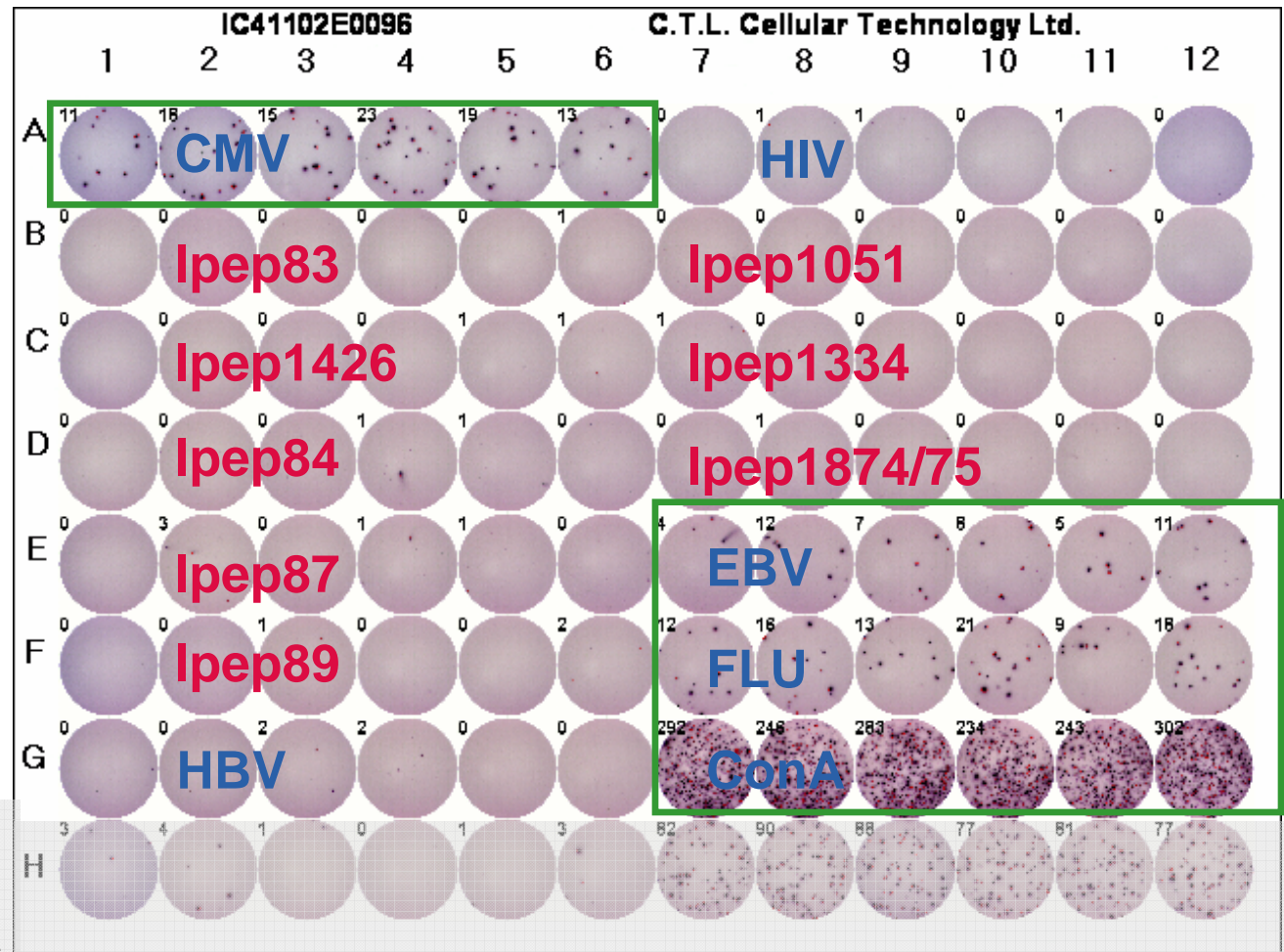
Klade et al. Gastroenterology 2008

* Study period: end 2002 - mid 2004
** Different dose levels

Interferon gamma ELISpot using frozen PBMC

ELISPOT: $\geq 3x$ OVER BACKGROUND, AT LEAST 15 PER MIO. PBMC

Positive Controls:
CMV, EBV,
Flu-peptides
Con A

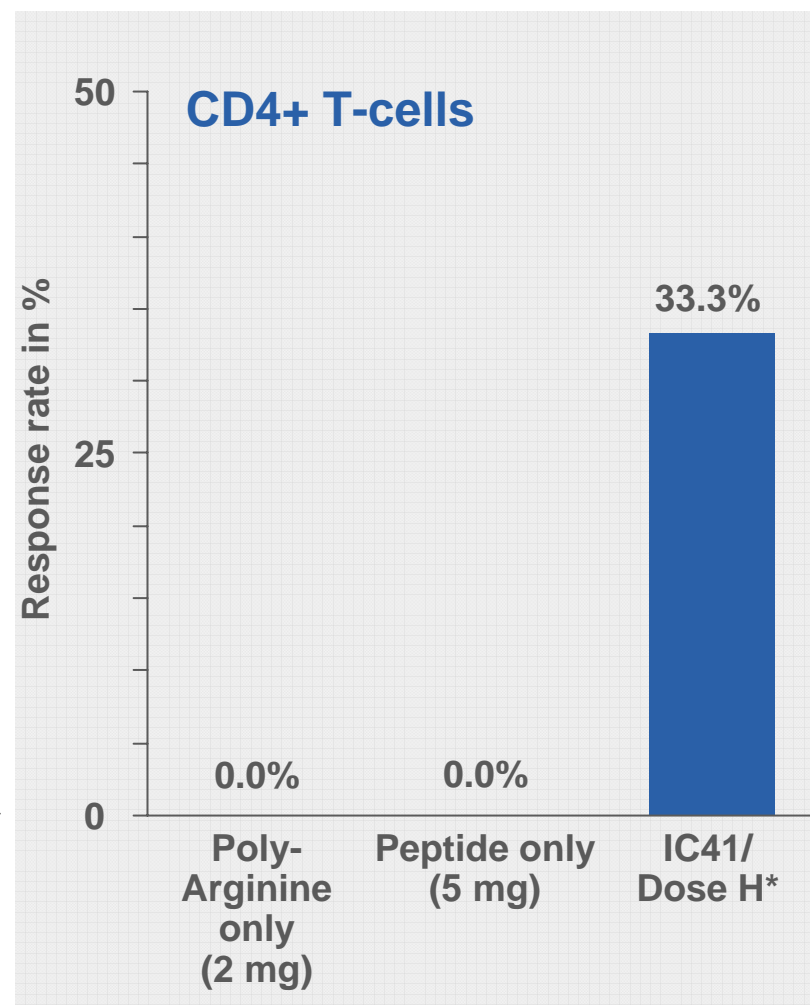
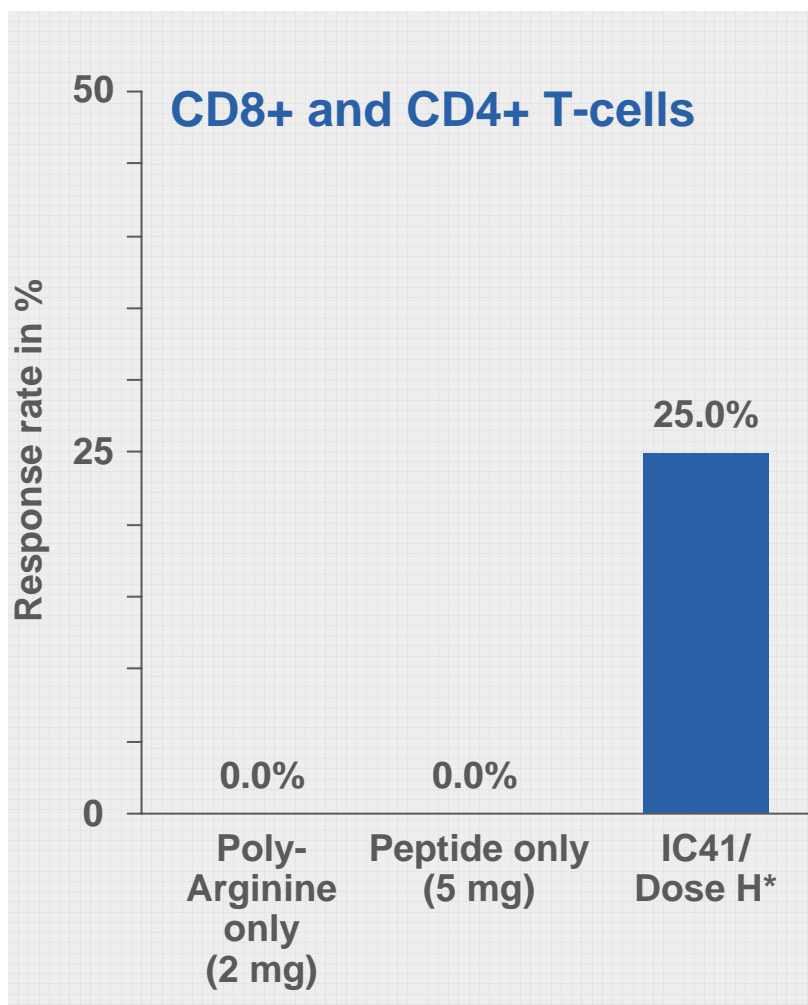


Assay standard:
control cells
HIV vs. CMV peptides

IC41 induces Th1/Tc1 type immune responses in non-responder patients

Phase II in non-responders

CLASS I AND II RESPONSE RATES (ELISPOT)



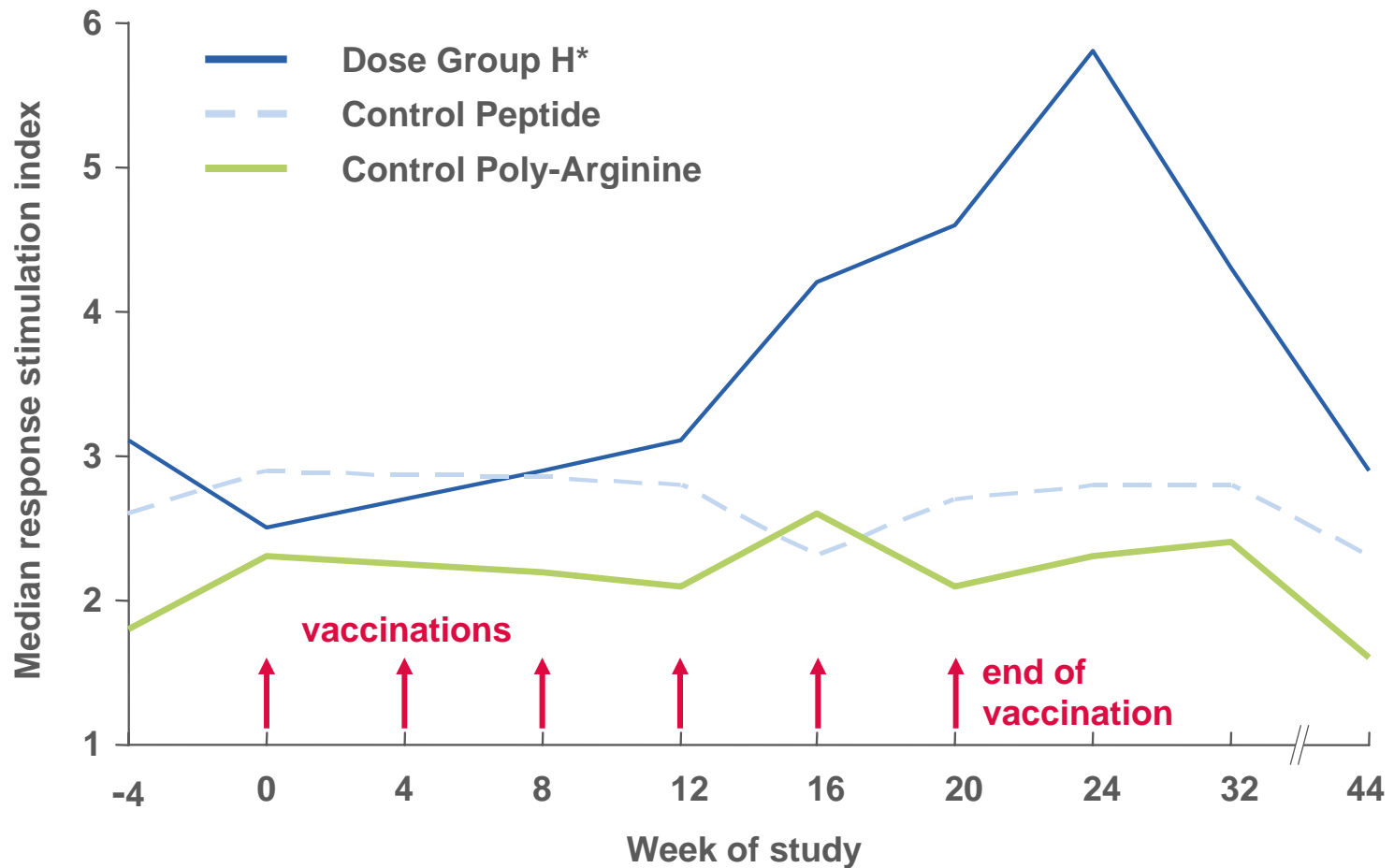
Klade et al.
Gastroenterology
2008,
Firbas et al., 2006

* 2.5 mg peptides;
2.0 mg Poly-Arginine

IC41 induces T-cell proliferation in non-responder patients

Phase II in non-responders

MEDIAN CLASS II T-CELL PROLIFERATION: DOSE GROUP H



Klade et al.
Gastroenterology
2008

* 2.5 mg peptides;
2.0 mg Poly-Arginine

Results of concluded Phase II study – IC41 already showed trend in efficacy

Phase II in non-responders

PHASE II NON RESPONDERS (IC41-1)

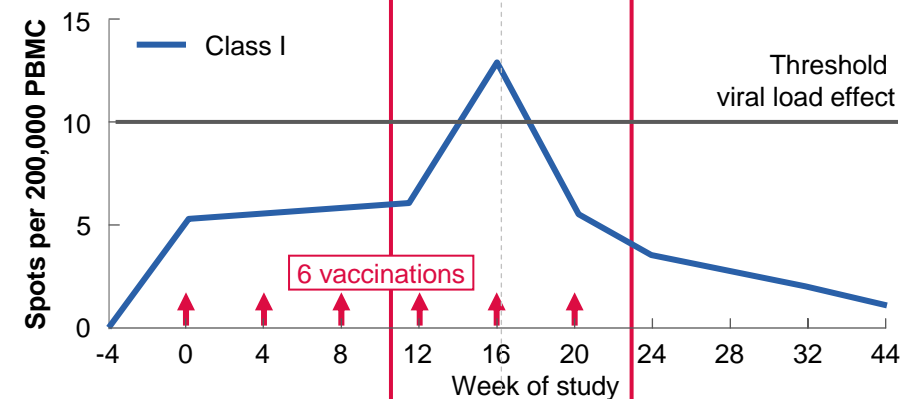
Group results of 1 Log responders in Phase II trial*

Group	Dosage	N	Resp.
K	5.00/2.00	2	17%
H	2.50/2.00	1	8%
G	2.50/1.25	0	0%
B	0.00/2.00	0	0%
C	5.00/0.00	0	0%

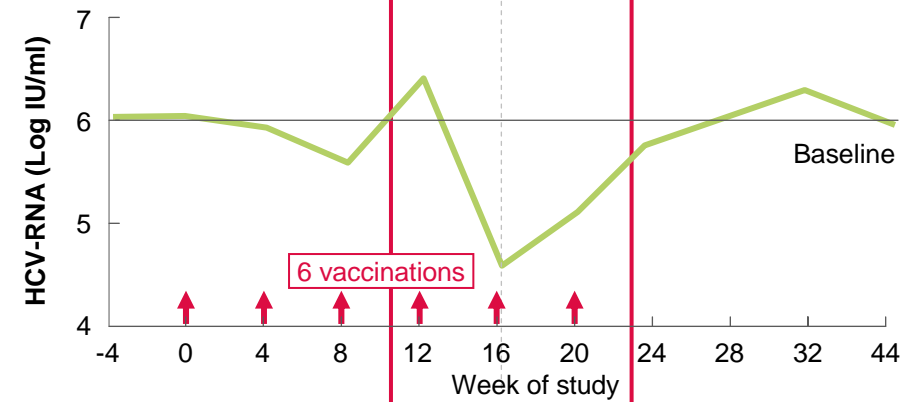
Class I responses of >10spots/200,000 are associated with transient viral load reductions

Results of patient with viral load reduction in high dose group*

ELISPOT



HCV-RNA

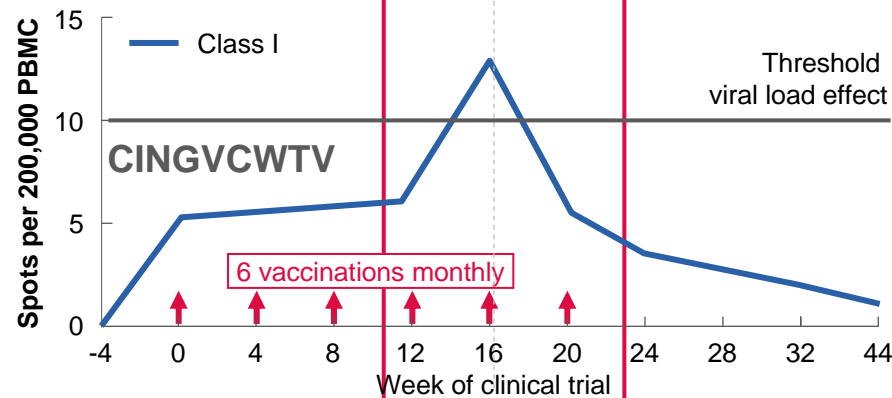


Evidence for mutational T-cell epitope escape in a patient responding to IC41-1 vaccination

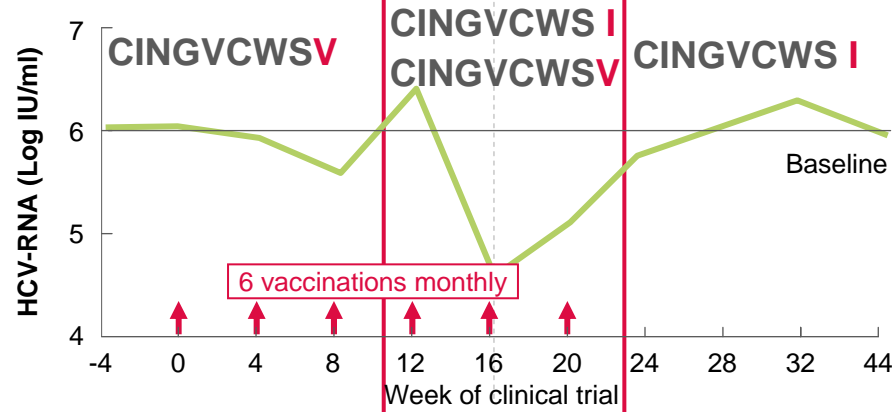
Phase II in non-responders

RESULTS OF PATIENT WITH VIRAL LOAD REDUCTION*

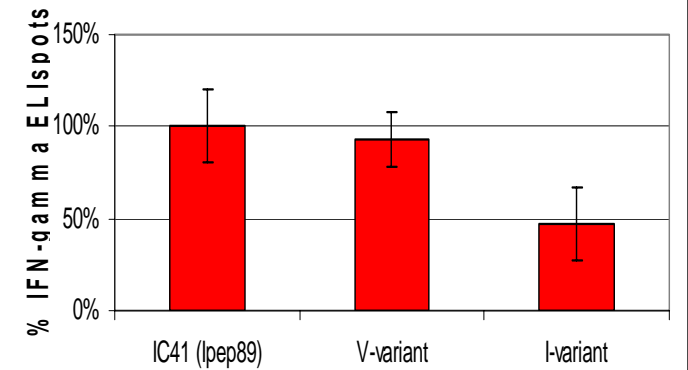
ELISPOT



HCV-RNA



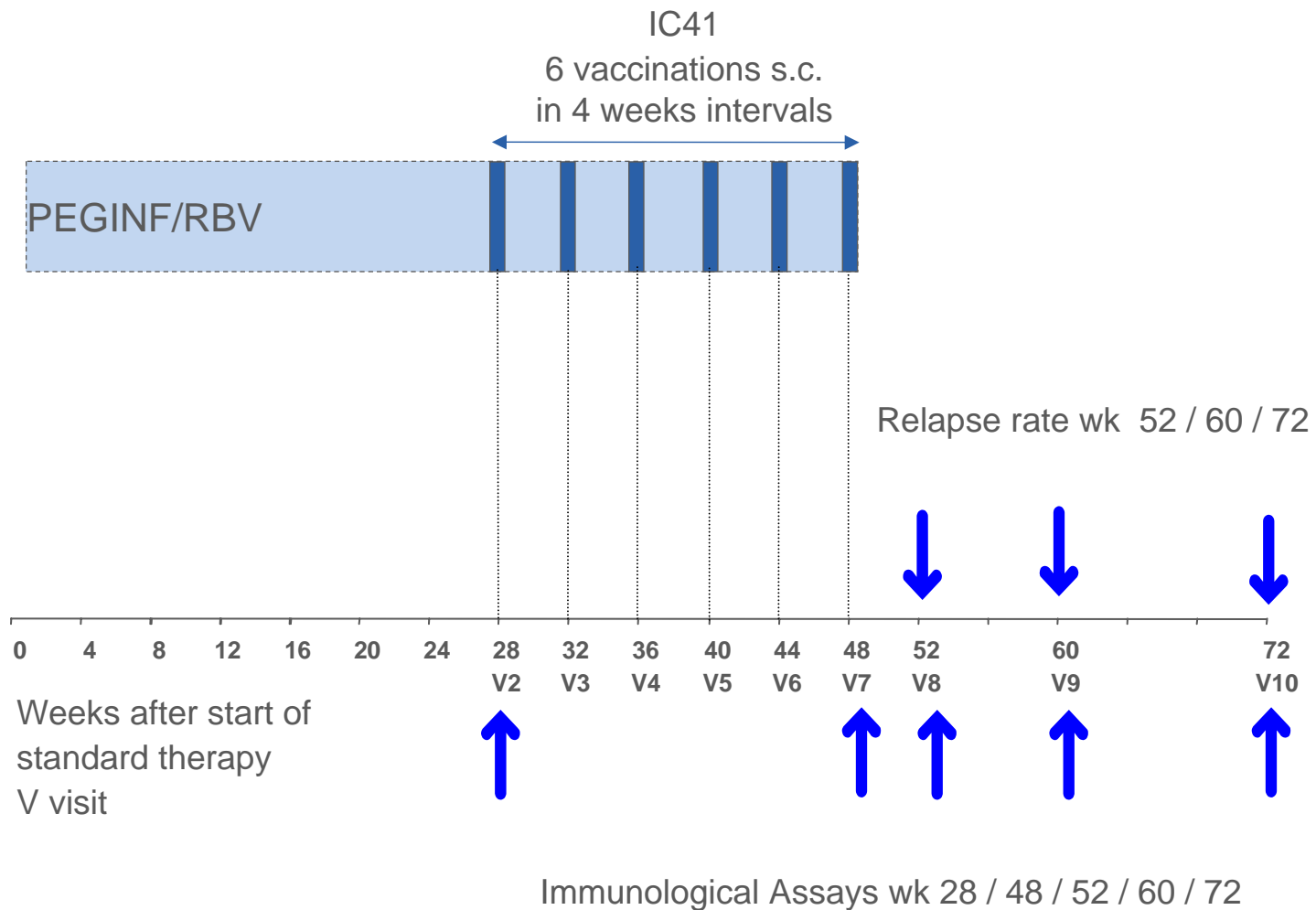
Impaired recognition of an HCV T cell epitope evolving in a single patient during vaccination



* Published and presented at the EASL Meeting in Vienna, April 2006

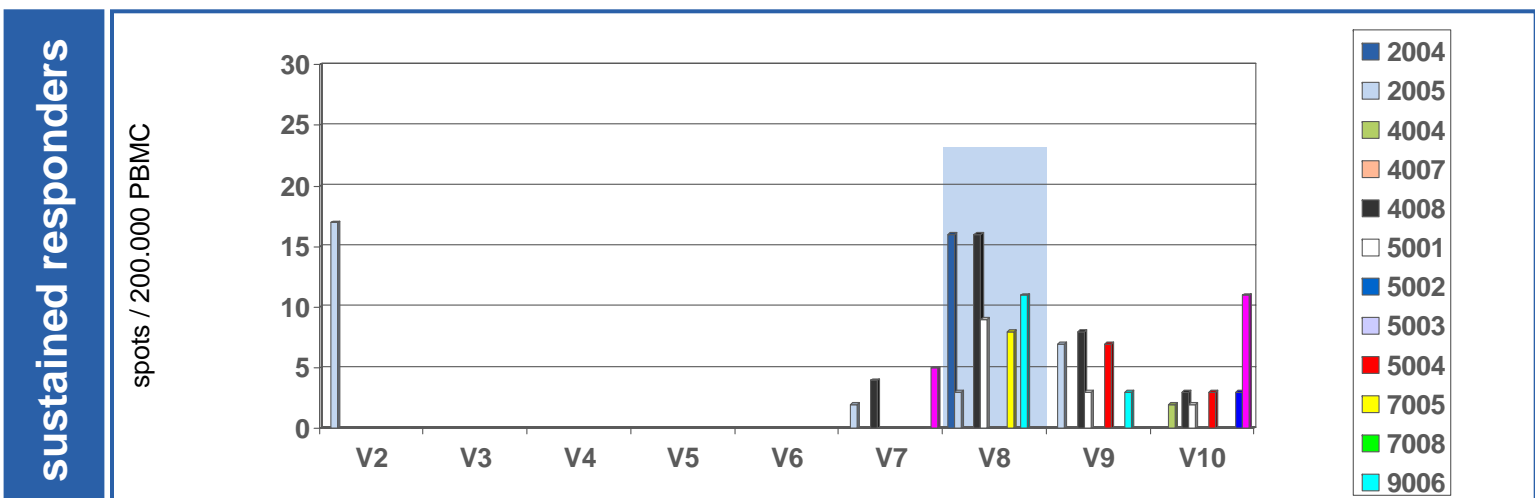
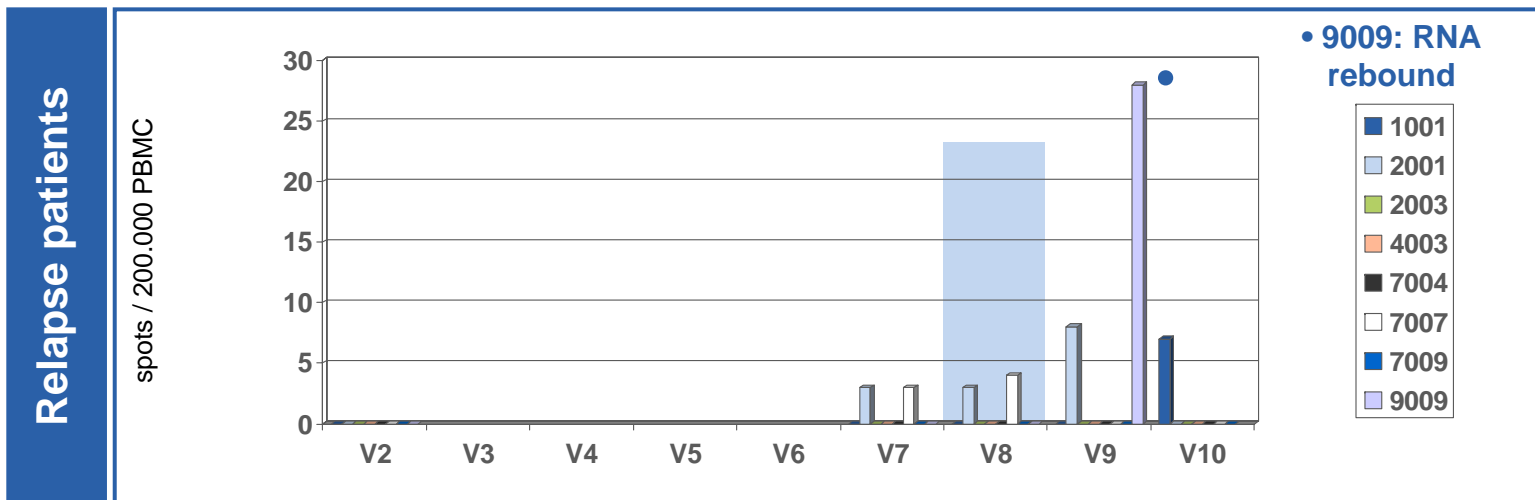
IC41-2: Combination with standard therapy

Patients with chronic hepatitis C of genotype I scheduled for standard treatment for 48 weeks already treated for 28 weeks and responded at week 12



Sustained responders show a stronger and more frequent T-cell response – Target Population*

INTERFERON γ ELISPOTS IN RELAPSED PATIENTS (N= 8) VS. SVR (N=14)



*Target Population N = 23, for 1 patient missing HCV-RNA data between V8–V10

Conclusions from non-responder patients (IC41-1) and late add-on to PEG-IFN/RBV (IC41-2)

- » favorable safety profile in chronic HCV patients with or without PEG-IFN/RBV standard therapy
 - » optimal vaccine dose (2,5 mg peptides / 2,0 mg poly-L-Arg)
 - » Th1/Tc1 type responses in chronic HCV patients, no apparent negative influence through PEG-IFN/RBV
 - » several transient 1 log Hepatitis C - RNA responders at optimal dose
 - » RNA responses associated with strongest CD8+ responses achieved
- » **T-cell immunogenicity requires optimization**
(rate, strength, breadth, sustainability)

Improving immunicity of IC41 in HLA-transgenic mouse model

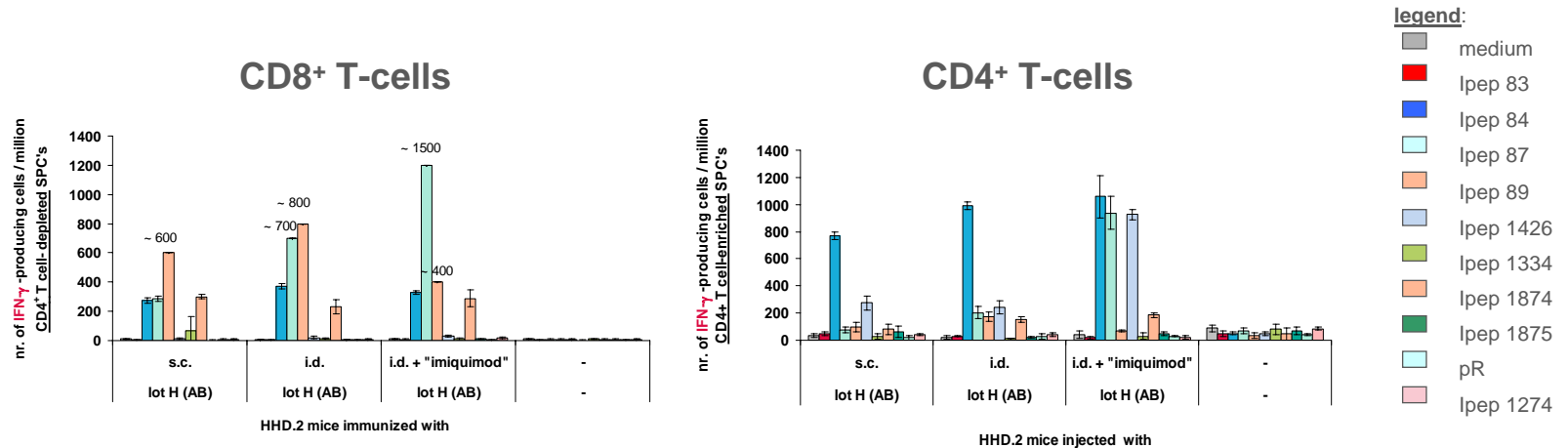
TEST APPLICATION SITES ± IMIQUIMOD

HHD.2 mice
Dose/100µl/mouse:
**100µg/peptide +
400µg pR**
(lot H in-house
mixture AB)

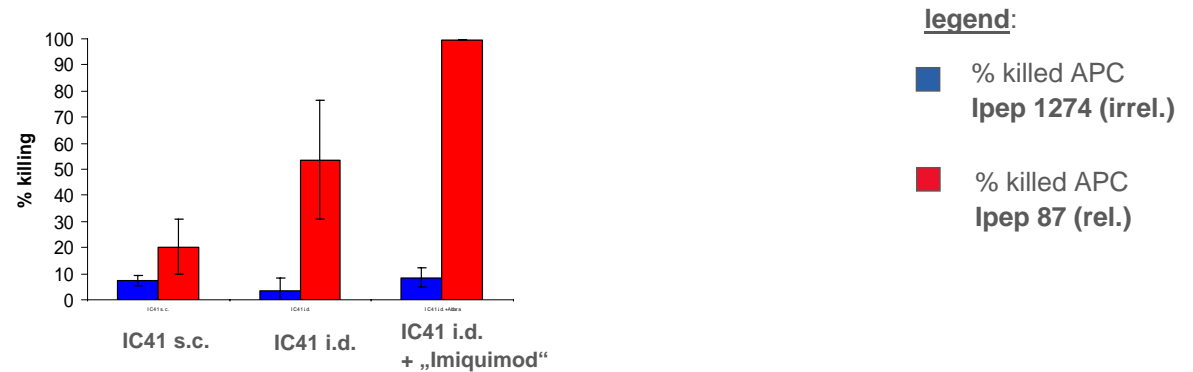
exp. scheme:
**day 0, 14, 28,
42, 56, 70**

s.c. or i.d.injection
day 7 after 6th inj.
IFN-g ELISpot
(spleen cells)

day 29 after 6th inj
APC transfer
day 30 after 6th inj
FACS analysis
spleen cells



In vivo CTL assay

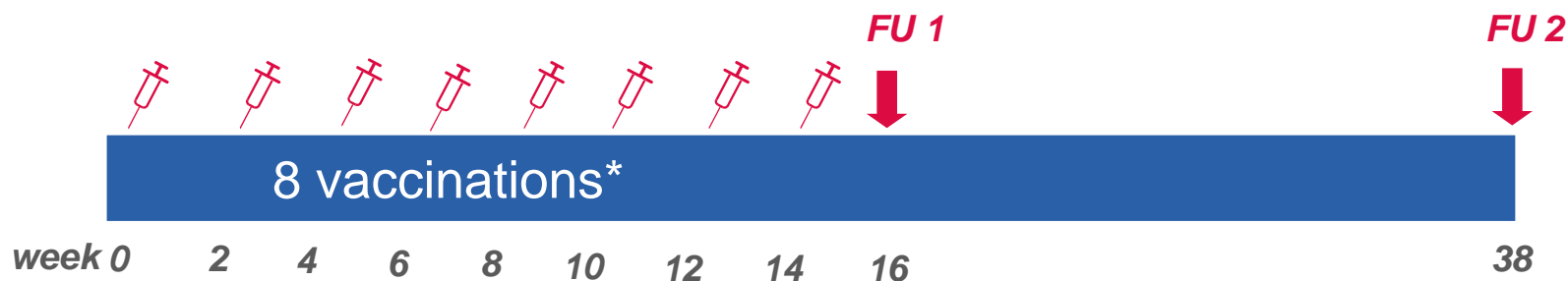


IC41-3 Study concluded January 2008

Phase II in
naïve
patients

OPTIMAL VACCINATION SCHEDULE IN TREATMENT NAIVE PATIENTS

- » 50 Chronic HCV patients, treatment naive, HCV Genotype 1.
Desired subset with low viral load at baseline



- » **First vaccination** on September 26 2006, first data Q2/2007

- » **Endpoints:**
 - Decline in HCV-RNA
 - T-cell response

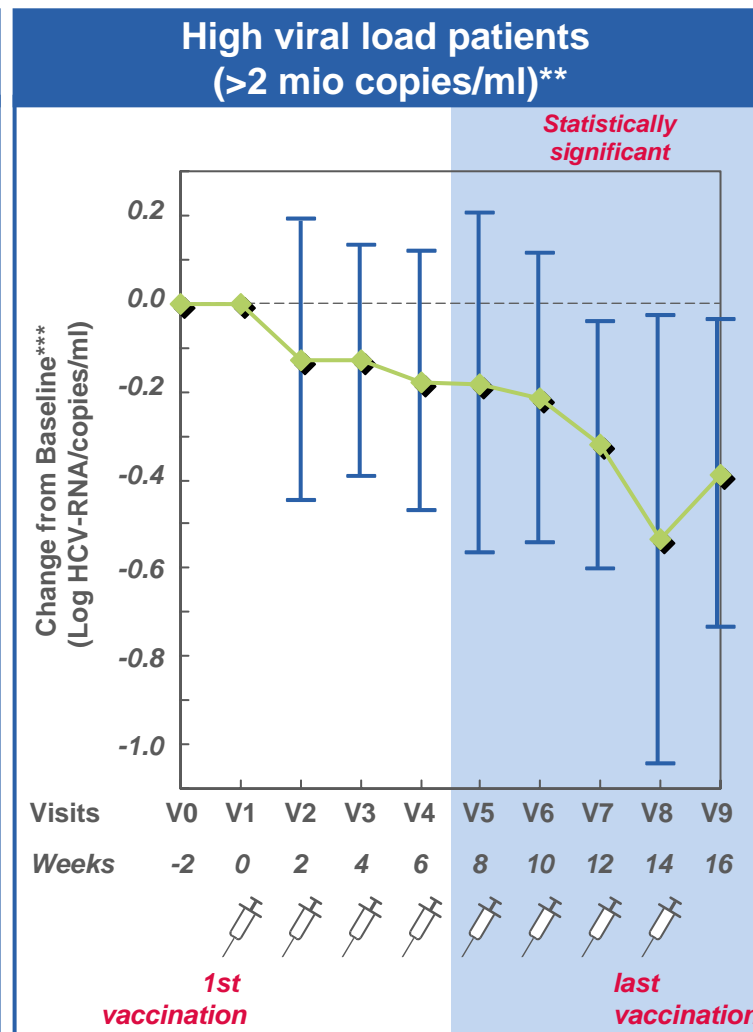
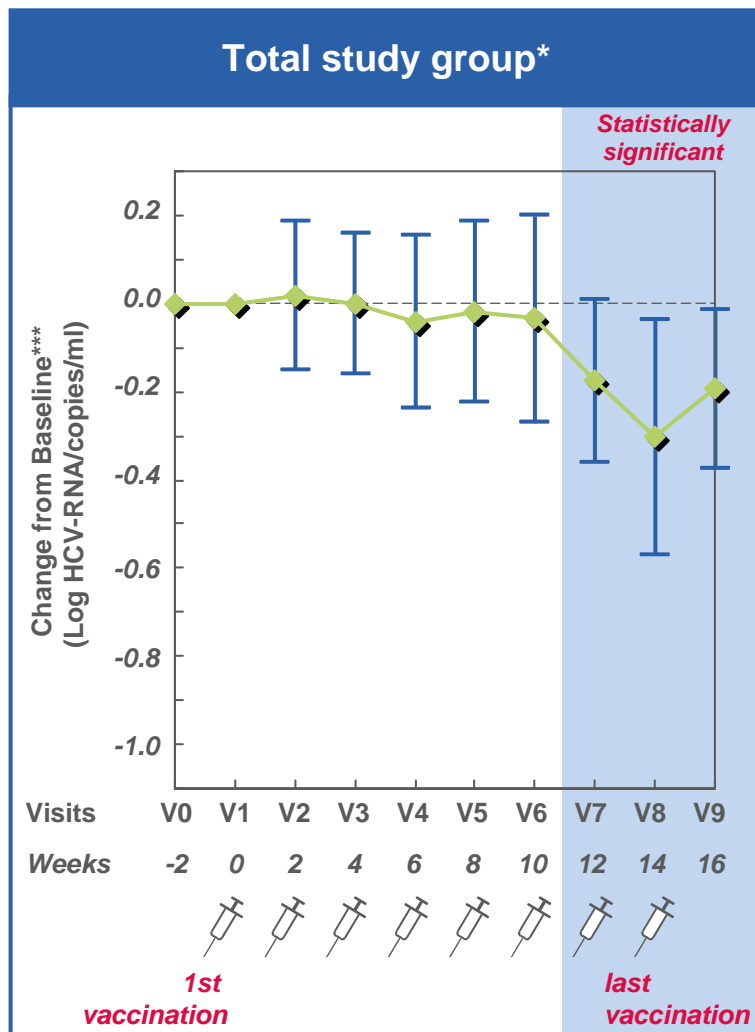
- » **Status**
 - Participating countries: Romania, Poland, Germany
 - End of recruitment on track for February 2007

* Bi-weekly;
intra-dermal;
topical Aldara®
(3M)

Primary endpoint met – a weak, but statistically significant HCV-RNA reduction

◆ Point Estimate
 Estimate

OVERVIEW IC41-3 PHASE II DATA



* 46 patients
 ** 25 patients
 *** 95% confidence intervals

Conclusions from IC41 trials

- » Favorable safety profile in chronic HCV patients with or without PEG-IFN/RBV standard therapy
- » Optimal vaccine dose / schedule identified
- » Th1/Tc1 type responses in chronic HCV patients, no apparent negative influence through PEG-IFN/RBV
- » Antiviral activity demonstrated in patients with strongest CD8+ responses, and treatment group with optimal vaccination

HCV therapeutic vaccination: Forward Strategy

Development of second generation vaccine

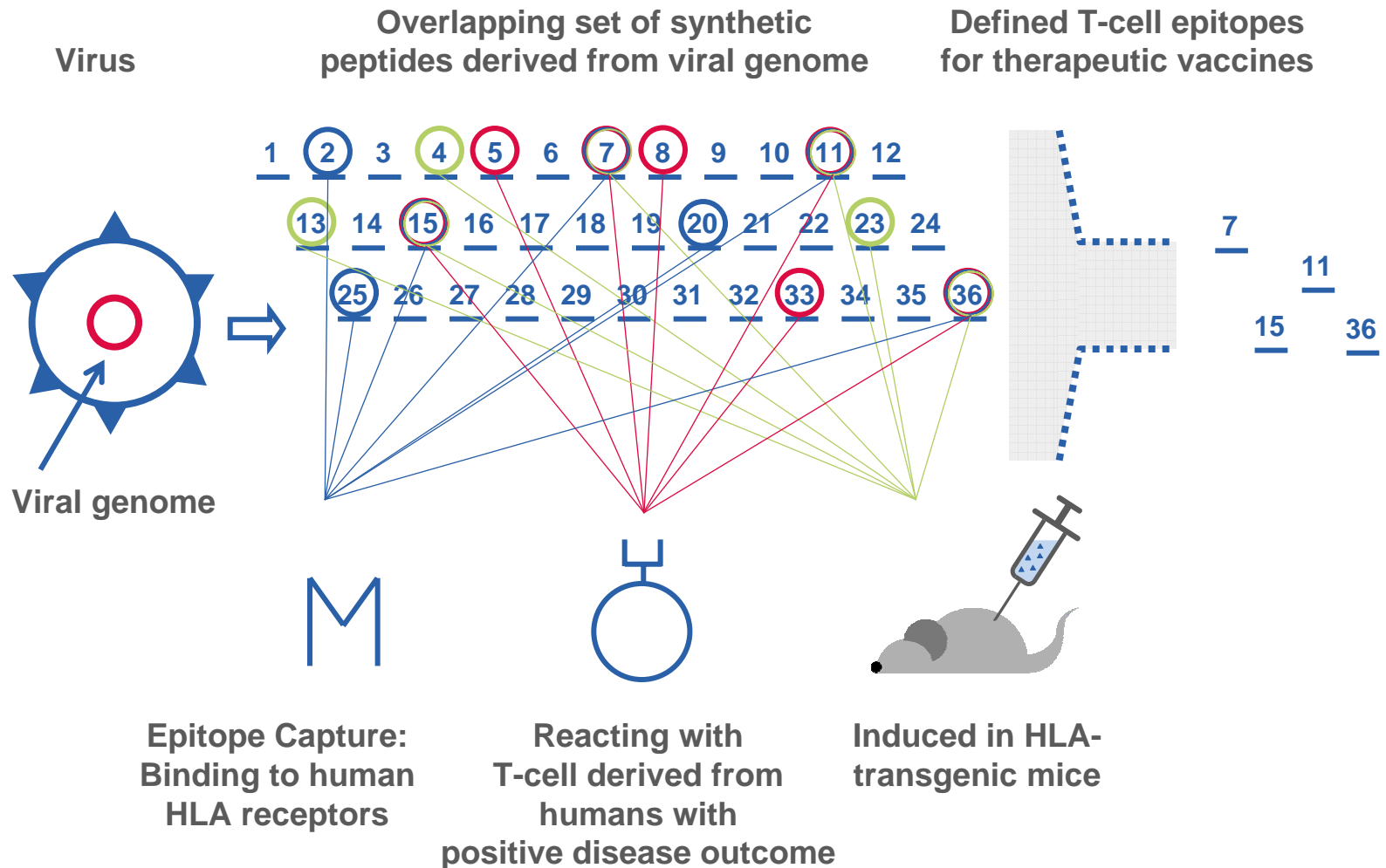
- » More & better peptides (HLA-restriction, efficacy)
- » Improved T-cell adjuvant (IC31[®])

Future plans: combination therapy

- » plus PEG-IFN/RBV
- » plus novel small molecule inhibitor

Identification of further T-cell peptides

T-CELL EPITOPE IDENTIFICATION PROGRAM



Reviewed in
Scharnagl & Klade
2007, Exp Rev
Vaccines 6:605-15.

Schalich & Klade
2008, Biol Chem

Kubitschke & Klade
2008,
in preparation

Identification of HCV vaccine candidate peptides beyond IC41

HLA-COVERAGE: 80-90% IN EUROPE, USA AND JAPAN

IVS: *in vitro* stimulation of PBMC from HLA-matched healthy donors

Peptide	Class I epitopes	Class II epitopes	Human PBMC screening	tg mice screening	Epitope Capture	Additional predicted epitopes
lpep 1835	A2, A3, B7	DR11	✓	✓ (B7 / lpep 1506)	+	
lpep 1829	A2, B7	DR1, 7, 11(?)	✓ (lpep1605, IVS)	✓ B7, (A2)	++(+)	A24
lpep 1799	B35	DR1, 4	✓	✓ (DR4 / lpep 1563)	++	
lpep 1798	A2, A3, A11	DR1, 4, 7	✓	(✓) (A2 no final data)	+++	A24
lpep 1827	A24	DR1, 7, 11	✓ (lpep1801)	Not applicable	+++	B8
lpep 1846	A2, A11(?), Cw7	DR1, 4, 7, 11	✓ (lpep1800, IVS)	✓ (DR4 / lpep 1650)	++++	A24
lpep 1547	A2	DR1, 4, 7, 11	(✓) (from Day et al.)	✓ DR4	++++	
lpep 1624	B60	DR7	✓	(as expected negative for A2, B7, DR4)	+	

PCT/EP2003/009482

Otava & Klade
AASLD 2004

Kubitschke & Klade
in preparation

IC31[®]: a TLR agonist comprising two chemically defined biodegradable components

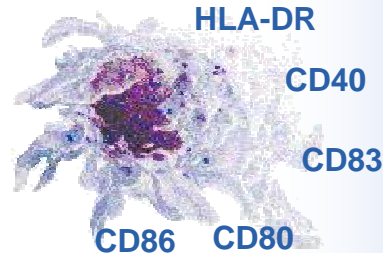
» KLK:

antimicrobial peptide H-KLKL₅KLK-OH

- Type 2 immune responses (+ proteins)
- Depot formation at injection site



- Enhancement of antigen and ODN1a uptake by APC

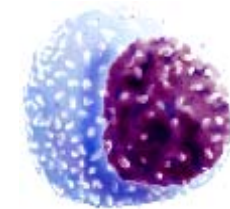


» ODN1a:

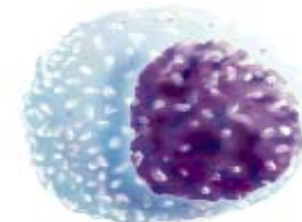
oligodeoxynucleotide oligo-(dIdC)₁₃ phosphodiester, ssDNA

- Type 1 induction
- Activation of APC (Dendritic Cells)
- TLR-9 / MyD88-dependent signaling

Potent and sustained
Th-1 / type 2
responses



T cell



B cell

IC31[®]: Induction of potent type 1 cellular immune responses

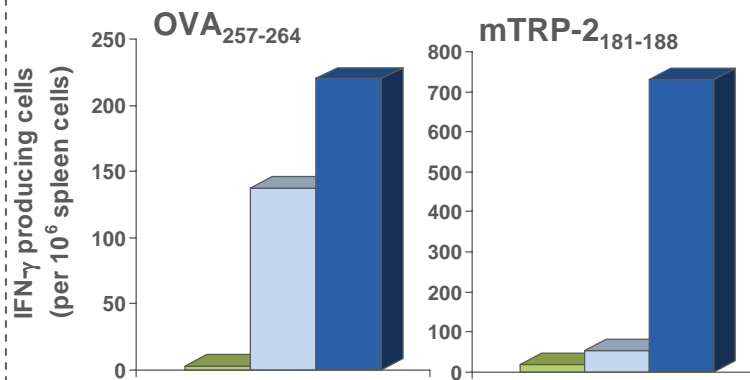
EXAMPLE: IMMUNIZATION WITH MODEL PEPTIDES

CTL - EFFECTOR CELLS

PEPTIDE-SPECIFIC IFN- γ PRODUCTION

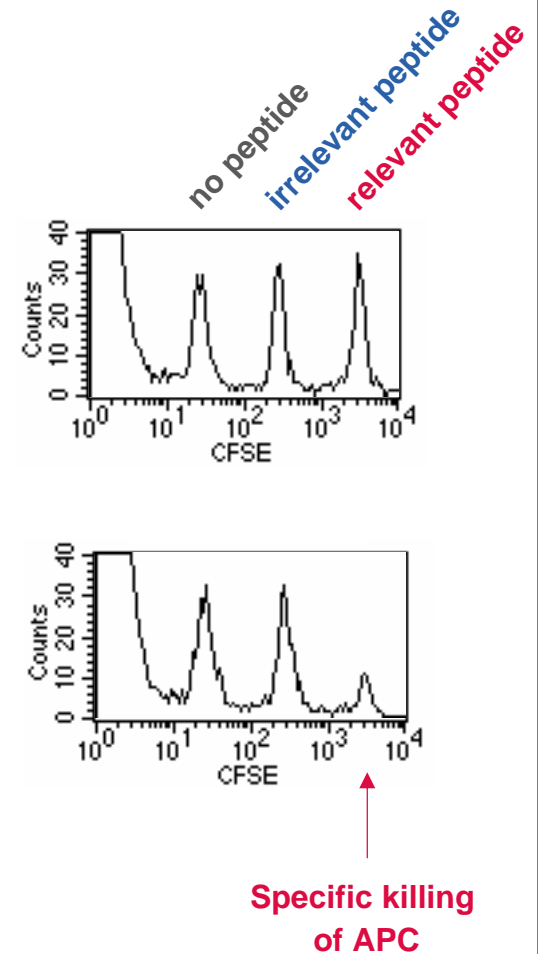
day 7 after single injection
IFN- γ ELISpot

- Alum
- CpG 1668
- IC31[®]



naive
or
mTRP-2₁₈₁₋₁₈₈

mTRP-2₁₈₁₋₁₈₈
+ IC31[®]



Protective immunity of a novel TB subunit vaccine adjuvanted with IC31[®]



STATENS
SERUM
INSTITUT

&



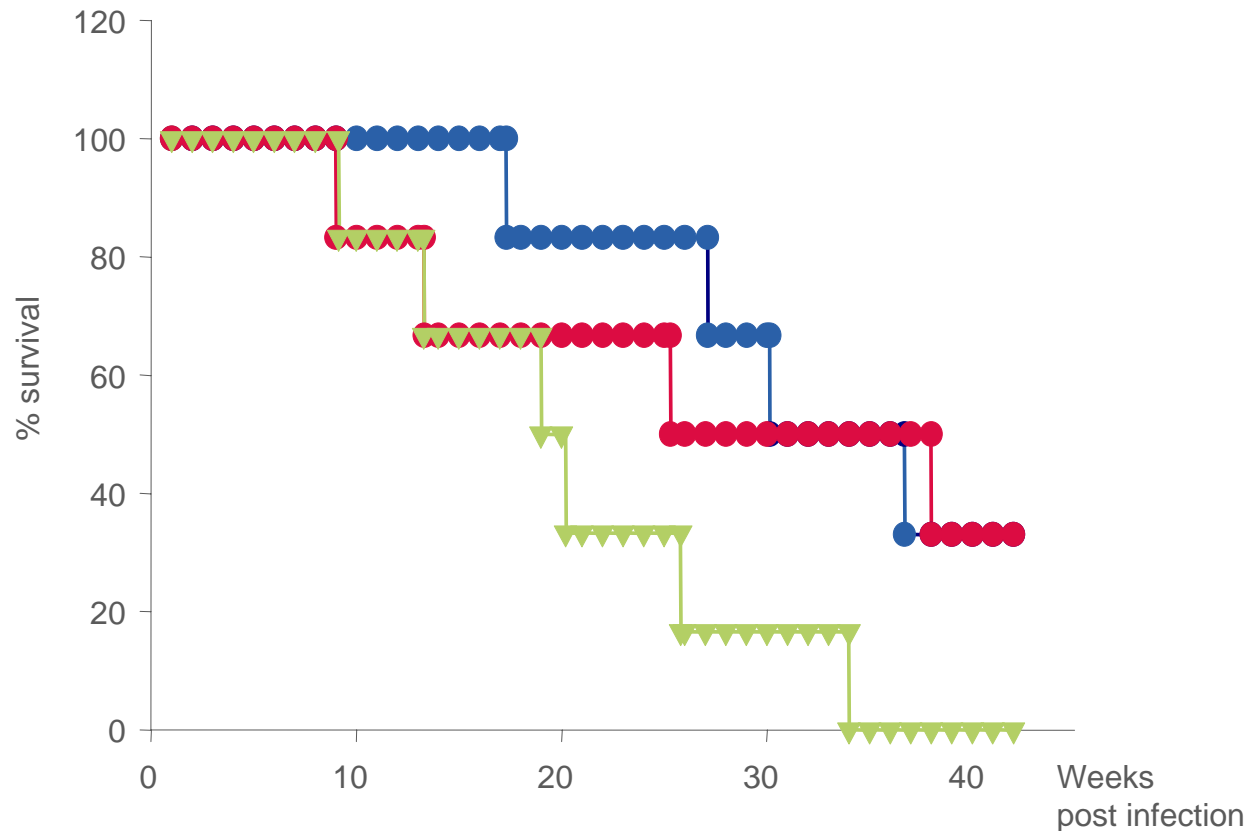
intercell
SMART VACCINES

- BCG
- Ag85B/
ESAT-6 +
IC31[®]
- ▼ Naive/
Saline

* 3x i.m.
injection, 4-
week interval

Aerosol
infection;
16 weeks after
first injection

PRECLINICAL EVALUATION – SURVIVAL (GUINEA PIG)*



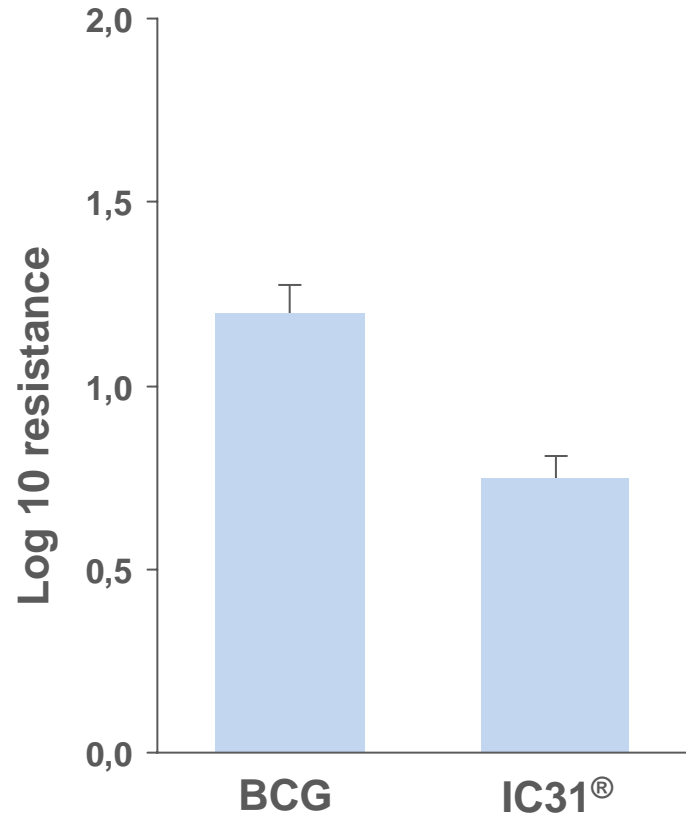
Protectivity is linked to IFN- γ producing T-cells indicative for Th-1 driven immunity



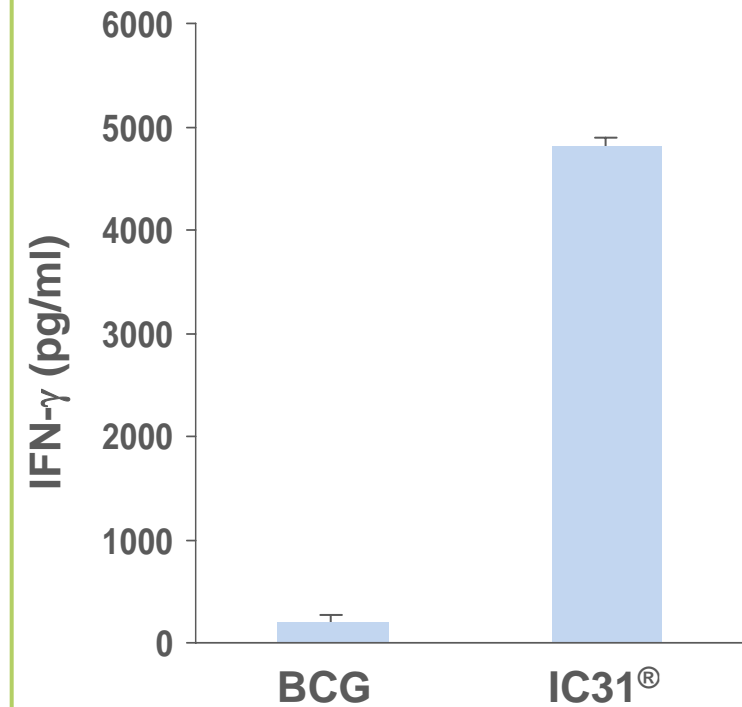
&

DEFINITION OF PROTECTION MARKERS (MOUSE MODEL)

RESIDUAL BACTERIA (lung)



IFN- γ production



Induction of antigen-specific T-cells in humans vaccinated with the novel TB subunit vaccines

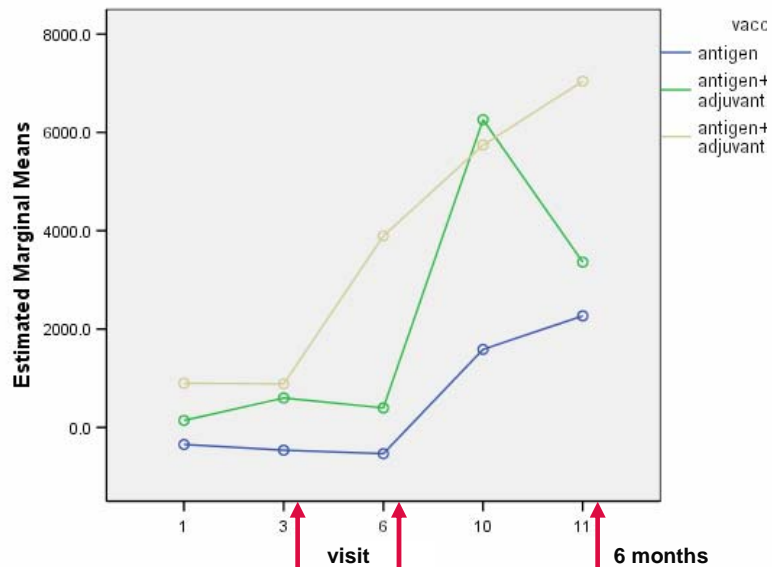


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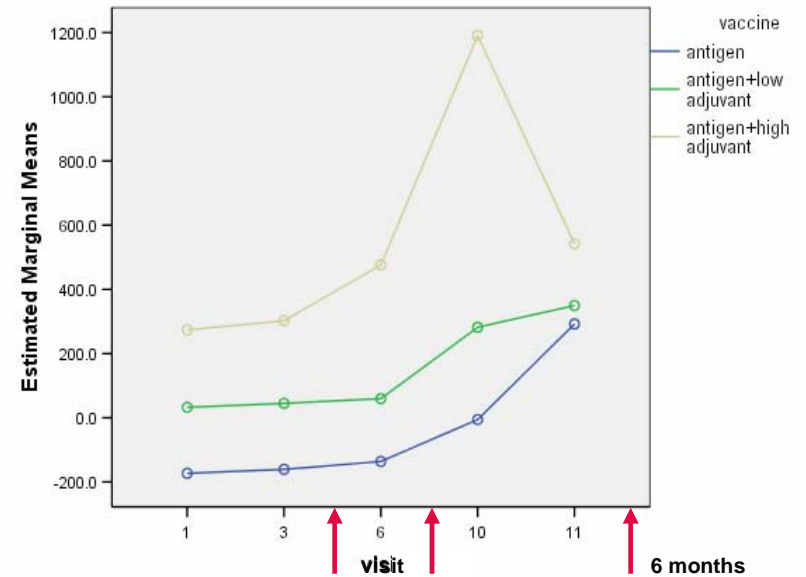


DATA FROM TB PHASE I STUDY: STRONG T_H-1 INDUCTION

IFN- γ in T-cell supernatants (Ag85B/ESAT-6-specific ELISA; Estimated Marginal Means)



Frequency of IFN- γ prod. T-cells (Ag85B/ESAT-6-specific ELISpot; Estimated Marginal Means)

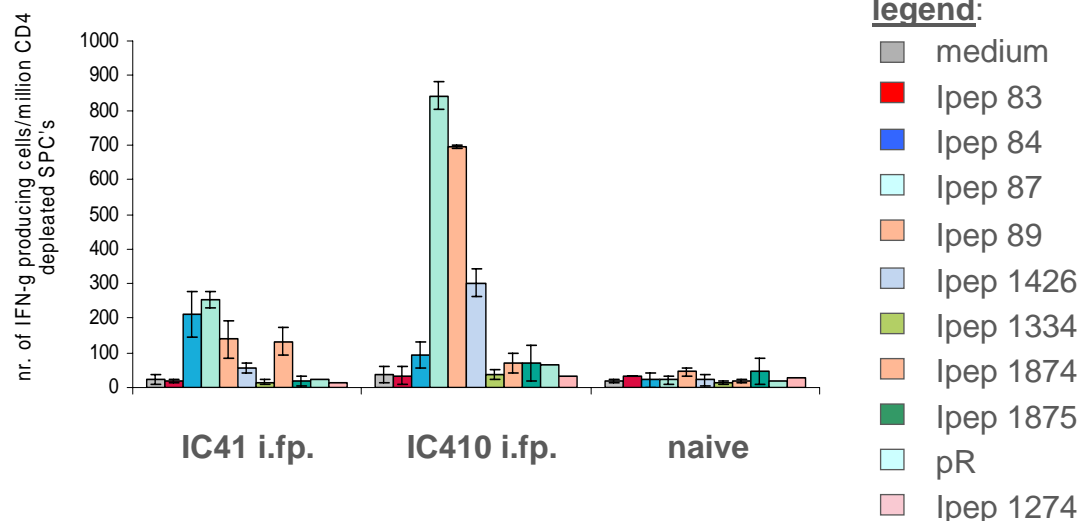


Dramatic improvement of IC41 by replacing poly(Arg) with IC31[®] (IC410)

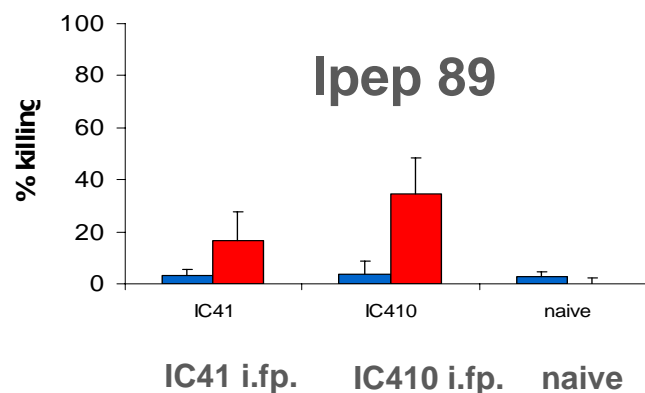
Dose/100µl/mouse:
IC41:
 200µg/peptide +
 400µg pR43
 (lot K, batch PD03126)

IC410:
 50µg/peptide +
 35nmol KLK+
 1.4nmol ODN1a
 (inhouse mixture)

IFN- γ PRODUCTION



CD8+ T CELL EFFECTOR FUNCTION



exp. scheme:
 day 0, 14, 28
 i. fp. injection
 day 34
 APC transfer
 day 35
 FACS analysis (LNC)
 ELIspot (spleen cells)

legend:

- % killed APC Ipep 1274 (irrel.)
- % killed APC Ipep 87, 89 (rel.)

Acknowledgments

INTERCELL

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HCV study group

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