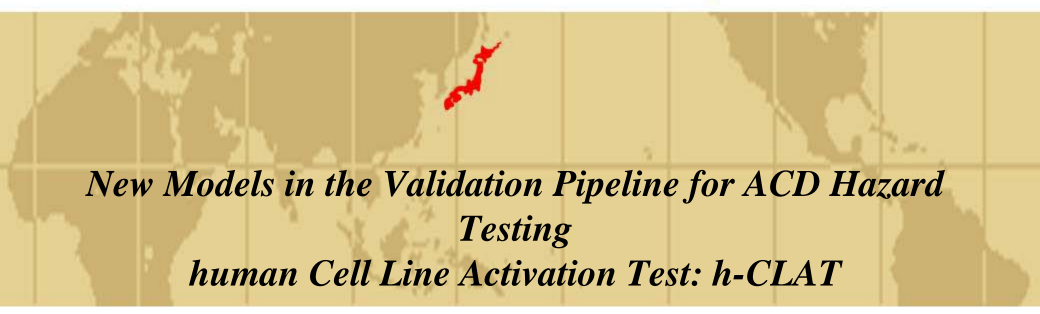


January 20, 2011

ICCVAM Workshop Series on Best Practices for Regulatory Safety Testing Assessing the Potential for Chemically Induced Allergic Contact Dermatitis

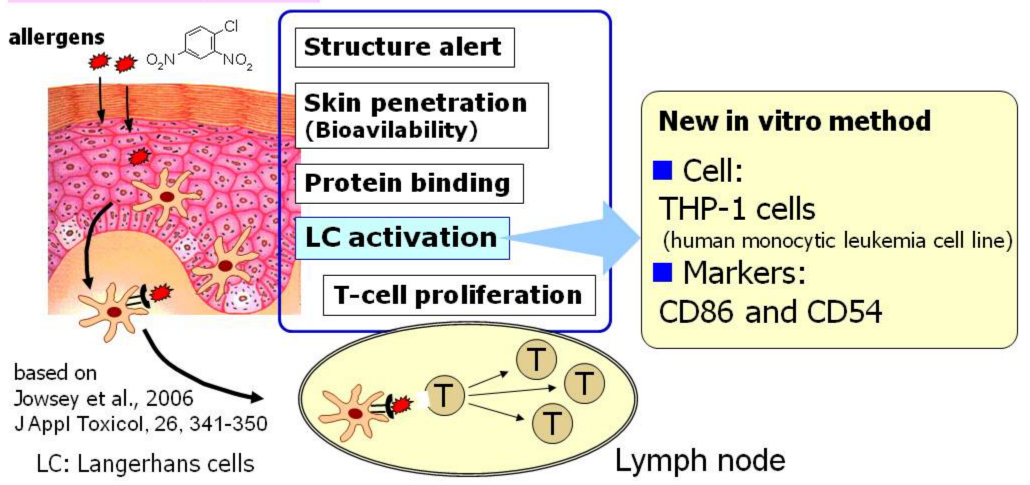


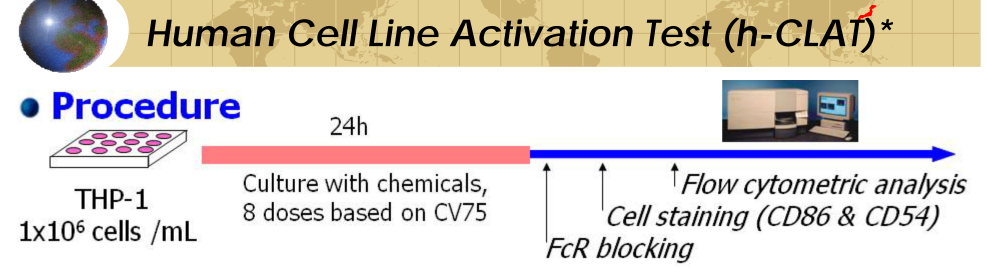
Hitoshi Sakaguchi, Kao Corporation Takao Ashikaga, Shiseido Co., Ltd

Approach for Developing of in vitro Methods

It is imperative to understand the mechanisms the sensitization (induction) phase of contact hypersensitivity (Vandebriel et al., 2005)

Induction phase





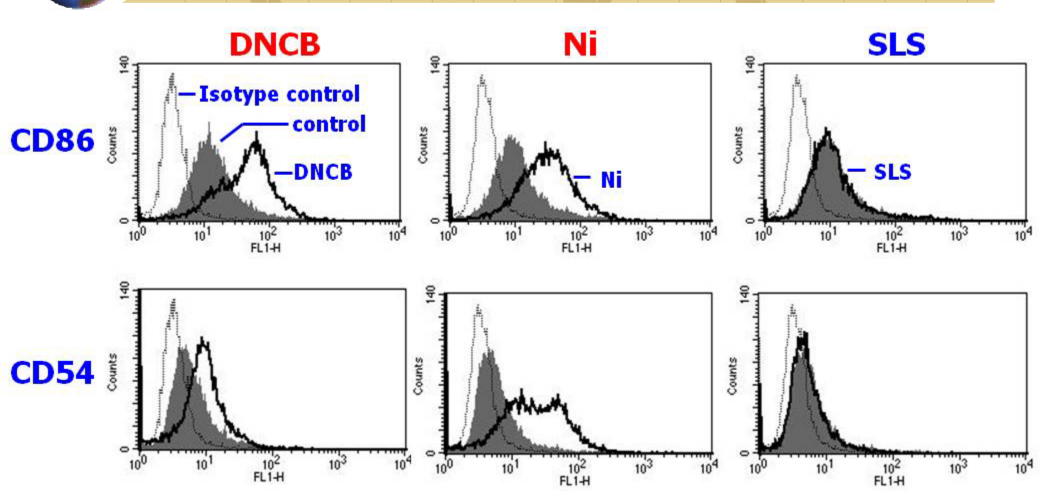
• Relative Fluorescence Intensity (RFI)



Prediction Model

- Viability \geq 50% by Propidium Iodide
- Positive criteria: CD86 RFI \geq 150% and/or CD54 RFI \geq 200%
- Positive: 2 of 3 independent data at any dose should exceed the positive criteria
- *: Ashikaga et al., 2006 Toxicol In Vitro 767-73., Sakaguchi et al., 2006 Toxicol In Vitro 774-84.

Histogram of CD86 / CD54 expression



DNCB and Ni (typical allergens) enhanced both CD86 and CD54 expressions but SLS (non-allergen) did not.

Miyazawa et al., Toxicology in Vitro 2007

Today's presentation

• Predictive capacity

- Evaluation of 117 chemicals by the h-CLAT to compare with LLNA
- Applicability domain
 - Applicability domain based on the data base
- Classification of skin sensitization potency
 - Using EC150 and EC200 values as the indicator
- Inter-laboratory study
 - Ring Trials in the COLIPA (5 labs) and Japan (7 labs)

Results of 117 test chemicals

	1 8/96 I.		1 No. 28, 103			H-EMIT				
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kaa ahme		0.000	P	10	+	23	271	- 3	100.0	DMASO
6(0MIDut:1.05)	10	0.005	P	TT -	+		2.21	1.1	12	Salina
androwski's base		0.00				+	16	2.5	23	DMISC
Sentoquintrat	11	0.0009	P	p	+	+	2.5	225	41	DMIST
die russyl acatomer	Extreme a	0.04	P			+		10.0	12.1	DMISC
		0.04	-			+	32	2.0	63	DMASO
etr ach lonn sail i cyl avil late A-Gint troich londo en ar nei ± DNCB		0.05	P	1		1.	23	200	SO SO	DMISC
Whenhoused home ide		0.05	-				0.95	0.90	18	DMISC
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lal ni a antrophridat	8	0.15	P	P		+	- 9 • ~~	2984	891.0	DMSD
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nru yî kramî de		8.2	P	- m 3		+	3.2	200	7.5	DMISC
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opyl gallate	Strong	0.12	P	. p		+	100	12.5	125.0	DMISO
thettchloutde		0.3	P	H	0.20	+	- 22	2.2	208.3	Salim
animphenit			P	11	+	1.1	0.00		8.0	DMASE
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		0.5	-		-	+		278.7	60.7	DMAST
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et hyblibe onso glutta con it rile			P	P	•	+		8.62	8.8	DMSD
ae upon at		12	P	- 26	12.00		- 8.÷	• • · ·	112.5	DMSD
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Phone 1, 2-program diana	19	13	Sec. 1		+	+	EL.5	52.7	1711	DMISC
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nyl pyridios		1.5		H	+		11.2	- 23	72.4	DMISO
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http://www.inconecust.9.7%)	10	1.9	P			-	8.23	7.60	24.7	Salira
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e thyl m ale are	1	5.8	P	- n	+	S			120.0	DMISO
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Phony for optional de hyde	10	6.3	12. 3			+	22	41.5	17.9	DMASE
Chloria anili ne		85	P	1		-	87.7		200.0	DMISO
rilablahyda		11			+	-	51.00		83	DMISC
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Nukada et al., WC7 2009, Ashikaga et al., ATLA 2010, Nukada et al., ESCD 2010

Comparative evaluation with LLNA and human

h-CLAT vs LLNA

Accuracy:

h-CL	.AT	VS	human	

		h-CLAT		
		+(83)	-(34)	
	+(85)	75	10	
LLNA	-(32)	8	24	

Sensitivity:	75/85(88%)
Specificity:	24/32(75%)
Positive predictivity:	75/83 (90%)
Negative predictivity:	24/34(71%)
	The second s

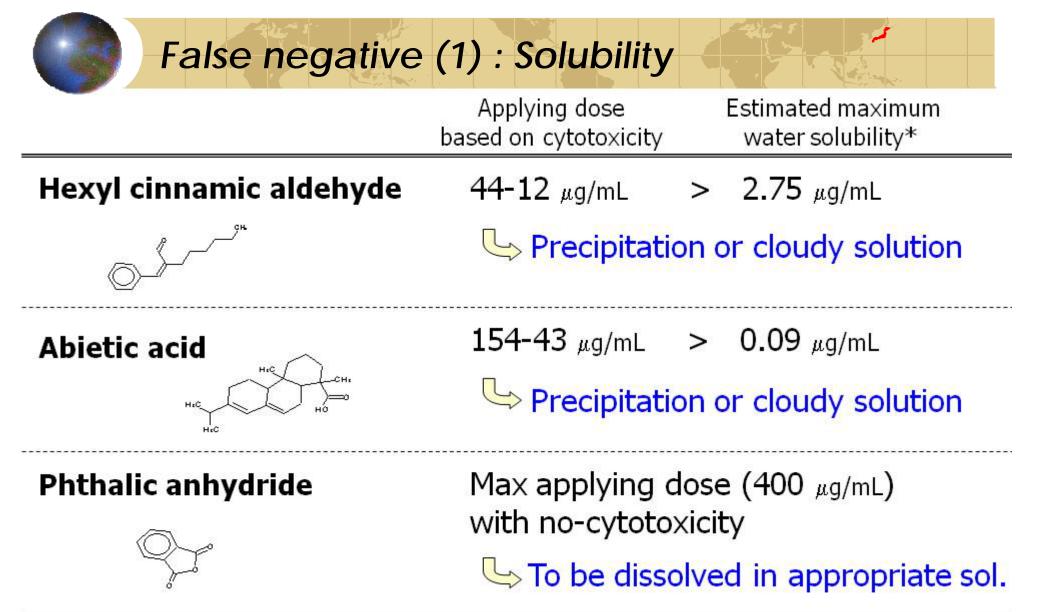
99/117 (85%)

		h-CLAT		
		+(51)	-(20)	
	+(55)	46	9	
Human	-(16)	5	11	

Accuracy:	57/71 (80%)
Negative predictivity:	11/20(55%)
Positive predictivity:	44/51(90%)
Specificity:	11/16 (69%)
Sensitivity:	46/55 (84%)

Good predictive capacity, but some false negative / positive

Nukada et al., WC7 2009, Ashikaga et al., ATLA 2010, Nukada et al., ESCD 2010

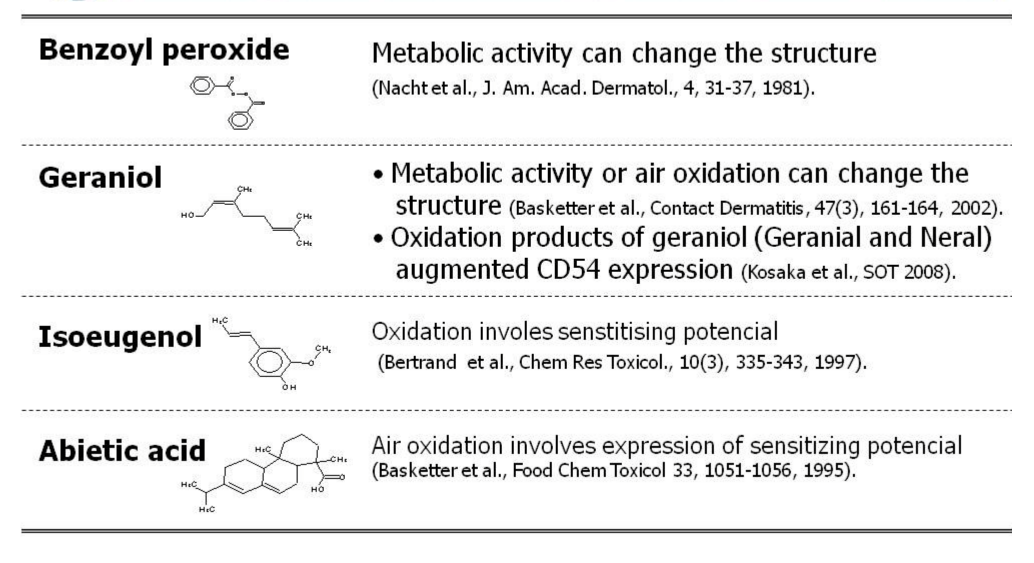


The chemical with poor water solubility is one of limitation

Ashikaga et al., ATLA 2010

*: Calculated with "Water frag" software.

False negative (2): Pro(Pre)-hapten



The h-CLAT had limitation for some pro- and pre-hapten

False negative (3) : Sensitivity

Weak sensitizers by LLNA classification

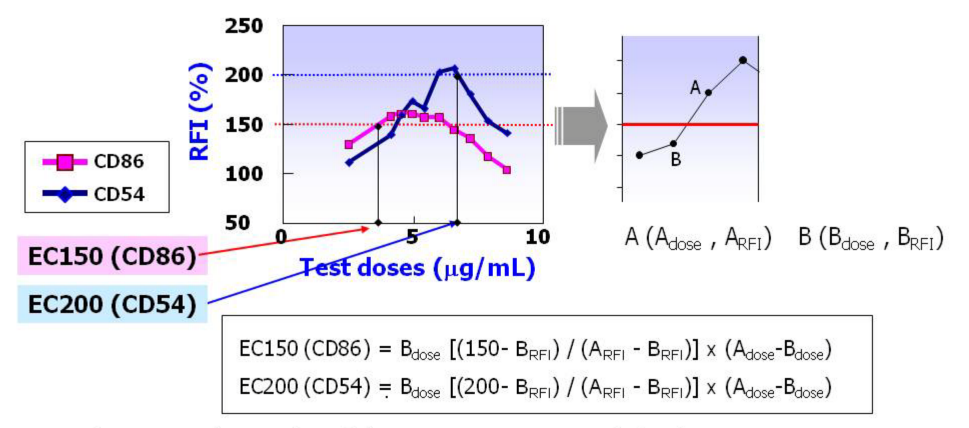
1-Bromohexa	ne Cyclamen	aldehyde	Butyl glycidyl ether		
CH₅ Br-			Åŧ		
LLNA class	Number of tested chemicals	Number of false negatives	Sensitivity (%)		
Extreme	8	0	100		
Strong	16	2	88		
Moderate	24	2	92		
Weak	23	5	78		

Several weak sensitizers could not enhance CD86/CD54 expression Ash

Ashikaga et al., ATLA 2010

Correlation between h-CLAT and in vivo data

EC150 / 200 (Estimated concentration of RFI 150 / 200)



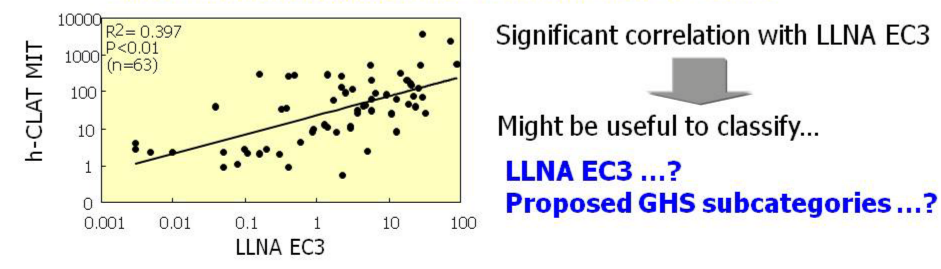
The intermediate value of three experiments was defined as EC150 or EC200.

Calculated based on the calculational procedure of LLNA EC3

Classification of skin sensitization potency

Minimum Induction Threshold of h-CLAT – MIT (h-CLAT) -

determined as a smaller value of either EC150 or EC200



Ref. Proposed GHS subcategories for skin sensitization based on LLNA EC3 and the example of prediction

Subcategory	Animal test results (using LLNA data)	Cut off (h-CLAT)	Accuracy(%)
1A (Strong sensitizer)	EC3 ≤ 2%	MIT 10 μg/mL	78.8
1B (Weak sensitizer)	EC3 > 2%		/0.0

COLIPA and Japanese Ring Trials

Purpose

- Protocol transferability
- Inter-laboratory reproducibility
- Predictive capacity

Goals

- Identify unexpected problems with either test design or procedures
 - Protocol optimization/standardization
- Identify problems with data analysis / interpretation
 - Prediction model refinement

Members

- COLIPA: P&G, L'Orel, Henkel-Phnion, Shiseido and Kao
- Japan: Kanebo Cosmetics, Kose, Lion, Nippon Menard Cosmetic, Pola Chemical Industries, Shiseido and Kao

COLIPA 4th Ring Trial summary data

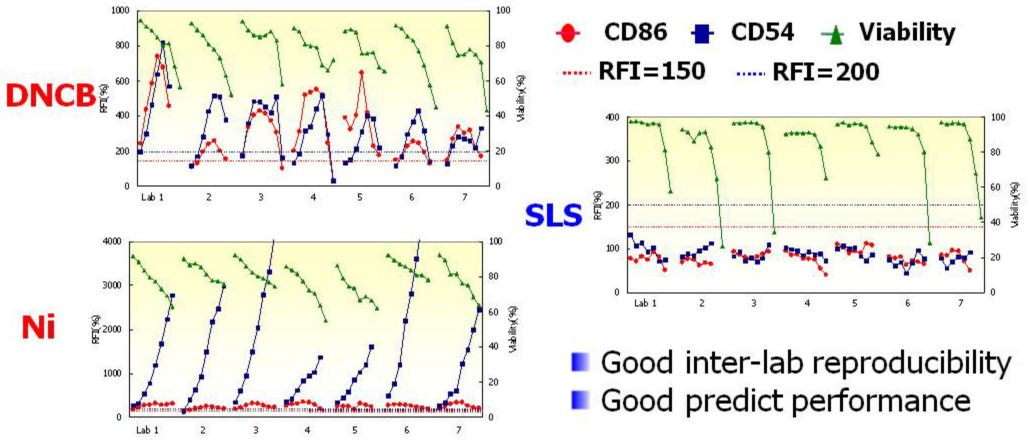
Chemical	Potency	Lab B	Lab C	Lab D	Lab E
PPD	Strong	+ (2/3)	+ (3/3)	+ (3/3)	+ (3/3)
Methyldibromo glutaronitrile	Strong	+ (3/3)	+ (3/3)	+ (2/3)	+ (3/3)
2-Mercaptobenzothiazole	Strong	+ (3/3)	+ (3/3)	+ (3/3)	+ (3/3)
Cinnamic Aldehyde	Moderate	- (1/3)	+ (3/3)	+ (2/3)	+ (3/3)
Tetramethylthiuram Disulfide	Moderate	+ (3/3)	+ (3/3)	+ (3/3)	+ (3/3)
Glycerol	NS	- (0/3)	- (0/3)	- (0/3)	- (0/3)
Salicylic Acid	NS	+ (3/3)	+ (3/3)	+ (2/3)	+ (3/3)

7 test chemicals (5 allergens, 2 non-allergens), 4 labs

- Cinnamic Aldehyde : one false negative data
- Salicylic acid : false positive in all labs
- Good inter-laboratory reproducibility
- Almost good predict performance

Japanese 1st Ring Trial

- 3 test chemicals (2 allergens, 1 non-allergen), 7 labs
- Test doses were same in all labs



Ashikaga et al., AATEX 2008

Summary

- Predictive capacity (117 chemicals)
 - Good prediction performance (accuracy: 85%/80% between the h-CLAT/human and LLNA) was observed.
- Applicability domain
 - Possible applicability domain was solubility, metabolic activity, sensitivity, etc.
- Classification of skin sensitization potency
 - MIT might be useful to predict the allergic potency of chemicals classified by GHS classification
- Inter-laboratory study
 - COLIPA : 15 chemicals, approx 85% predicted correctly
 - Japan : 8 chemicals, approx 96% predicted correctly
 - Good inter-lab reproducibility and predictive performance

ECVAM prevalidation study

• Liaison:

• JaCVAM and ICCVAM

Test methods:

- Direct Peptide Reactivity Assay (DPRA)
- Myeloid U937 Skin Sensitization Test (MUSST)
- human Cell Line Activation Test (h-CLAT)
- Main purpose
 - · The assessment of the robustness and reliability
- Experimental design
- 24 coded chemicals in three (or four) laboratories each for the assessment of the within- and between-laboratory reproducibility

Acknowledgments

Shiseido Co., Ltd Sakiko Sono Makie Ishikawa Katsurako Yoneyama Morihiko Hirota Shigenobu Hagino Hiroshi Itagaki Kao Corporation Masaaki Miyazawa Yuuko Nukada Nanae Kosaka Yukiko Terada Yuuichi Ito Hiroyuki Suzuki

Thank you for your attention