

#### 4. Validation Management structure

This validation study was performed by the Japanese Society for the Alternative to Animal Experiments (JSAAE).

The management structure of the study is shown in Figure 1.

##### 4-1. Validation Management Group

The Validation Management Team (VMT), which plays a central role overseeing the conduct of the validation study, includes:

- 1) Goal statement
- 2) Project plan including objective
- 3) Study protocol / amendments
- 4) Outcome of QC audits
- 5) Test chemicals
- 6) Data management procedures
- 7) Timeline/ study progression
- 8) Study interpretation and conclusions
- 9) Reports and publication

The final decision on which laboratories participate in the validation study is the responsibility of the VMT.

Members:

A chair (Hajime Kojima, JaCVAM: Japanese Centre for the Validation of Alternative Methods)  
The sponsor representative: representatives of JSAAE (Takashi Omori; Kyoto Univ., Kenji Idehara; Daicel Chemical Co. and Isao Yoshimura; Tokyo University of Science)  
The sponsor representative, LabCyte EPI-MODEL24suppliers and lead lab (Masakazu Kato : Japan Tissue Engineering Co., Ltd, J-TEC)

##### 4-2. Chemical selection, acquisition, coding and distribution

- 1) Definition of selection criteria
- 2) *Chemical selection*
- 3) *Liaise with suppliers*
- 4) *Final check of chemicals provided*
- 5) *Acquisition*
- 6) *Coding*
- 7) *Distribution*

Member

Hajime Kojima, JaCVAM

##### 4-3. Independent biostatisticians

- 1) Approve spreadsheets
- 2) Collect data
- 3) Analyse data

Members:

Takashi Omori: Kyoto Univ., Etsuyoshi Mlyaoaka and Kenya Ishiyama: Tokyo University of Science

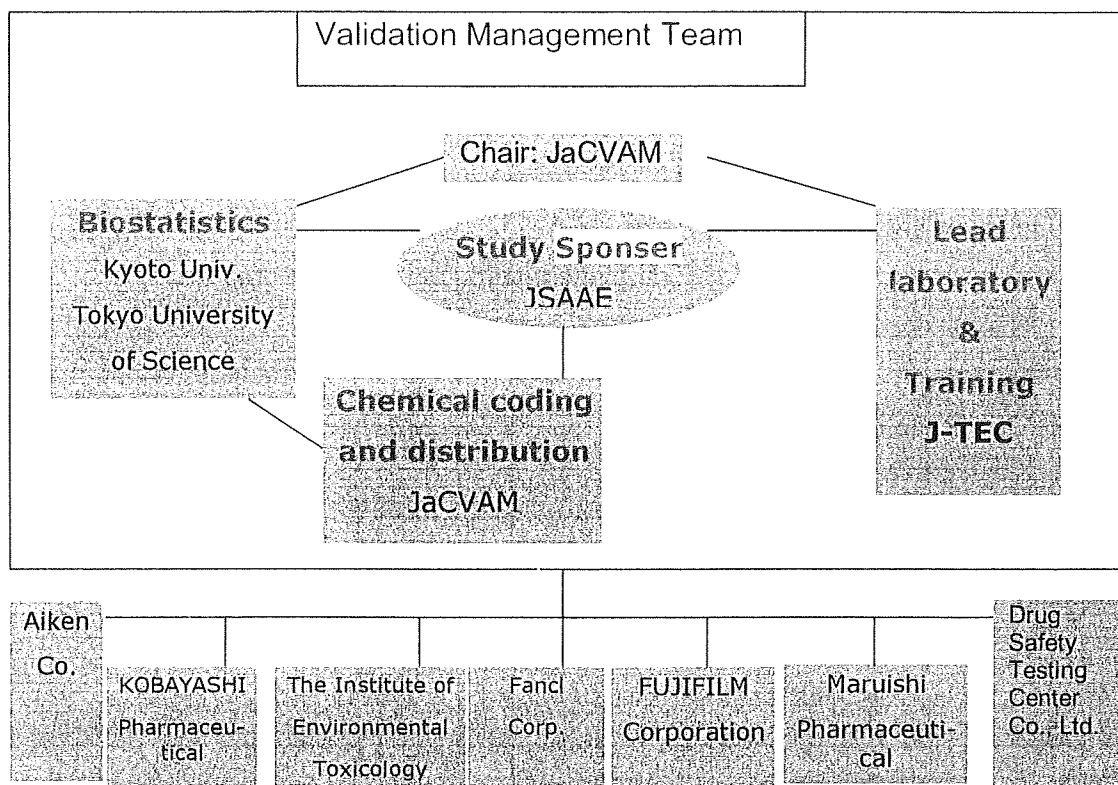


Figure 1. Management structure of the JSAAE skin irritation validation study

#### 4-4. Participating laboratories

The laboratories participating in the study are to be defined as shown in Fig. 1.

The following 6 laboratories participated in the validation study for the evaluation of the LabCyte EPI-MODEL24 assays:

- Laboratory 1 – Aiken Co., Ltd. (Yoko Ando and Yui Asako)
- Laboratory 2 – KOBAYASHI Pharmaceutical Co., Ltd. (Yoshihiro Yamaguchi and Maki Nakamura)
- Laboratory 3 – The Institute of Environmental Toxicology (Tadashi Kosaka and Koichi hayashi)
- Laboratory 4 – Fancl Corp. (Tamie Suzuki and Runa Izumi)
- Laboratory 5 – FUJIFILM Corporation (Atsuko Yuasa, and Shinichi Akimoto)  
This laboratory was not participated at the 3<sup>rd</sup> Phase study.
- Laboratory 6 – Maruishi Pharmaceutical Co., Ltd. (Yukihiko Watanabe and Osamu Mitani)
- Laboratory 7 – Drug Safety Testing Center Co., Ltd. (Shinsuke Shinoda and Saori Hagiwara)

A lead laboratory is also identified as J-TEC (Mr. Masakazu Kato and Mr Toshihiro Yokouchi). This laboratory was not participated in the validation study.

Each laboratory also was responsible for complying with GLP-like principles and specifying QA aspects.

#### 4-5. Sponsorship

The study was managed and finance by JSAAE and J-TEC .

##### 1)JSAAE finance:

- the management of the study (VMT meetings)
- the independent statistical support (biostatistician)
- the responsible for the chemicals purchase, coding and distribution to the laboratories
- the independent QC audit of the data
- the publication of the study

##### 2)J-TEC finance:

- the lead laboratories for the test method
- training for the participating laboratories
- the independent QC audit on the LabCyte EPI-MODEL24
- the financial assistance for the participated laboratories

#### 5. Study design

Before this validation study, the training course using LabCyte EPI-MODEL24 was performed by J-TEC on April, 2008. All technicians from each laboratory participated at this training course.

Three phases of validation studies were performed. In the 1<sup>st</sup> phase, we confirmed the transferability of the test protocol and assessed its reproducibility, by testing three coded chemicals (ethanol, glycerol and naphthalen acetic acid) and a positive control (5% sodium lauryl sulfate solution) in seven laboratories between June and July of 2008.

In the 2<sup>nd</sup> phase study, we confirmed the intra- and inter-laboratory reproducibility robustness, and the correlation of test using 19 new chemicals tested in reference to the original EPISKIN performance standards (ECVAM, 2007: Appendix 5) . These tests were conducted by 7 laboratories between September 2008 and January of 2009.

In the 3<sup>rd</sup> phase study, we confirmed the intra- and inter-laboratory reproducibility robustness, and the correlation of test using 6 chemicals tested in reference to the new EPISKIN performance standards (ESAC statement, 2009: Appendix 3). This study was conducted by 6 laboratories, which attend the 1<sup>st</sup> and 2<sup>nd</sup> phase validation study between April to May, 2009.

#### 6. Test Chemical

##### 6-1. Chemicals Selection and list

In 1<sup>st</sup> phase study, JaCVAM selected three coded chemicals (ethanol, glycerol and naphthalen acetic acid) to test.

According to the original ESAC Performance Standard (ESAC statement,2007) in 2<sup>nd</sup> Phase, the VMT selected 19 new chemicals to test in Table 1. One chemical, tri-isobutyl phosphate (No. 13) on the chemical list reference for the original ECVAM Performance Standard cannot be purchased on the Japanese market. The VMT is responsible for the final approval of the chemicals proposed by JaCVAM. To avoid any potential bias in the final selection, the laboratory representatives on the VMT were not party to these discussions, nor were they informed of the final list of test chemicals for either phase of the validation study.

According to the new ECVAM performance standard (ESAC statement, 2009) in 3<sup>rd</sup> phase, the VMT selected 6 new chemicals tested in Table 2. The final approval of the chemicals proposed by JaCVAM is the responsibility of the VMT. To avoid any potential for bias in the final selection, the laboratory representatives on the VMT did not be party to these discussions, nor were they made aware of the chemicals finally approved for testing in either phase of the validation study.

Table 1. Reference test chemicals and codes

No.	Chemical	CAS number	GHS label	In vivo score (PII)	Laboratory						
					a	b	c	d	e	f	g
01	1-bromo-4-chlorobutane	6940-78-9	no	0	A-01	B-099	C-077	D-115	E-133	F-031	G-049
02	diethyl phthalate	84-66-2	no	0	A-02	B-100	C-078	D-116	E-134	F-032	G-050
03	di-propylene glycol	25265-71-8	no	0	A-03	B-081	C-079	D-117	E-135	F-033	G-051
04	naphtalen acetic acid	86-87-3	no	0	A-04	B-082	C-080	D-118	E-136	F-034	G-052
05	allyl phenoxy-acetate	7493-74-5	no	0.3	A-05	B-083	C-061	D-119	E-137	F-035	G-053
06	isopropanol	67-63-0	no	0.3	A-06	B-084	C-062	D-120	E-138	F-036	G-054
07	4-methyl-thio-benzaldehyde	3446-89-7	no	1	A-07	B-085	C-063	D-101	E-139	F-037	G-055
08	methyl stearate	112-61-8	no	1	A-08	B-086	C-064	D-102	E-140	F-038	G-056
09	allyl heptanoate	142-19-8	no	1.7	A-09	B-087	C-065	D-103	E-121	F-039	G-057
10	heptyl butyrate	5870-93-9	no	1.7	A-10	B-088	C-066	D-104	E-122	F-040	G-058
11	hexyl salicylate	6259-76-3	no	2	A-11	B-089	C-067	D-105	E-123	F-021	G-059
12	terpinyl acetate	80-26-2	no	2	A-12	B-090	C-068	D-106	E-124	F-022	G-060
13	5(W/V %) SLS				A-13	B-091	C-069	D-107	E-125	F-023	G-041
14	1-decanol	112-30-1	Category 2	2.3	A-14	B-092	C-070	D-108	E-126	F-024	G-042
15	cyclamen aldehyde	103-95-7	Category 2	2.3	A-15	B-093	C-071	D-109	E-127	F-025	G-043
16	1-bromohexane	111-25-1	Category 2	2.7	A-16	B-094	C-072	D-110	E-128	F-026	G-044
17	$\alpha$ -terpineol	98-55-5	Category 2	2.7	A-17	B-095	C-073	D-111	E-129	F-027	G-045
18	di-n-propyl disulphide	629-19-6	Category 2	3	A-18	B-096	C-074	D-112	E-130	F-028	G-046
19	butyl methacrylate	97-88-1	Category 2	3	A-19	B-097	C-075	D-113	E-131	F-029	G-047
20	heptanal	111-71-7	Category 2	4	A-20	B-098	C-076	D-114	E-132	F-030	G-048

1) CAS No.: Chemical abstracts service registry number.

2) PII: Primary irritation index.

Table 2. Test chemicals and code.

No.	Chemical	CAS number	GHS label	In vivo Score	Laboratory					
					a	b	c	d	f	g
A	Cinnamaldehyde	104-55-2	no	2	A-151	B-176	C-196	D-216	F-236	G-256
B	2-Chloromethyl-3,5-dimethyl-4-methoxypyridine HCl	322-76821	Category 2	2.7	A-154	B-173	C-192	D-211	F-233	G-253
C	Potassium hydroxide (5%aq)	168-21815	Category 2	3	A-156	B-175	C-194	D-213	F-232	G-251
D	Benzenethiol, 5-(1,1-dimethylethyl)-2-methyl	7340-90-1	Category 2	3.3	A-153	B-172	C-191	D-214	F-234	G-254
E	1-Methyl-3-phenyl-1-piperazine	5271-27-2	Category 2	3.3	A-152	B-171	C-195	D-215	F-235	G-255
F	1,1,1-Torichloroethane	200-02463	Category 2	4	A-155	B-174	C-193	D-212	F-231	G-252

1)CAS No.: Chemical abstracts service registry number.

## 6-2. Deficit chemical

In Table 1, tri-isobutyl phosphate (No. 13) could not be used in the examination because it was not available in Japan. Therefore, a 5% SLS solution was used instead of tri-isobutyl phosphate. The data obtained with the 5% SLS solution were not used for calculating the predictivity of the test.

## 6-3. Chemical Coding and distribution

Independent coding and distribution of chemicals were contracted out by JaCVAM to an independent laboratory. The (company's name) is certified according to ISO 9001, EN 4500 and GLP, and has proven experience of reliable services. The codes were provided by JaCVAM.

## 7. Protocol

### 7-1. Protocol of the skin irritation test with LabCyte EPI-MODEL

In 2<sup>nd</sup> phase study, we used the SOP (ver. 5.0) and we used the SOP (ver. 6.1) in 3<sup>rd</sup> phase study. The revised points, which make the deletion measurement of IL-1 $\alpha$ , revise calculating formula of viability, classification used median of 3 trials and how to treat of volatile substances were shown in change tracking of the SOP (ver. 6.1: Appendix 6). The VMT made judgments that these revised points were minor and difference with the SOP (ver. 5.0) used by 2<sup>nd</sup> phase study and this version was little in the VMT meeting on July 17, 2009.

LabCyte EPI-MODEL tissues were shipped from the supplier on Mondays and delivered to recipients on Tuesdays. Upon receipt, the tissues were aseptically removed from the transport agarose medium, transferred into 24-well plates (BD Biosciences, CA, USA) with the assay medium (0.5 mL), and incubated overnight (37°C, 5% CO<sub>2</sub> humidified atmosphere). On the following day, the tissues were topically exposed to the test chemicals. Liquids (25  $\mu$ L) were applied with a micropipette, and solids (25 mg) were applied from microtubes and moistened with 25  $\mu$ L sterile water. If necessary, the mixture was gently spread over the surface of the epidermis with a microspatula. Viscous liquids were applied using a cell-saver-type tip with a micropipette. Each test chemical was applied to three tissues. In addition, three tissues serving as negative controls were treated with 25  $\mu$ L distilled water, and three tissues serving as positive controls were exposed to 5% SLS. After a 15-minute exposure, each tissue was carefully washed with PBS (Invitrogen, CA, USA) 10 times using a washing bottle to remove any remaining test chemical from the surface. The blotted tissues were then transferred to new 24-well plates containing 1 mL of fresh assay medium.

The treated and control tissues were incubated for 42 hours (37°C, 5% CO<sub>2</sub> humidified atmosphere). When the 42-hour post-incubation period was complete, blotted tissues were transferred to new 24-well plates containing 0.5 mL of freshly prepared MTT medium (1 mg/mL; Dojindo Co., Kumamoto, Japan) for the MTT assay and conditioned medium was collected to determine the interleukin-1 alpha (IL-1 $\alpha$ ) levels. Tissues were incubated for three hours (37°C, 5% CO<sub>2</sub> humidified atmosphere) and then transferred to microtubes containing 0.3 mL isopropanol, which completely immersed the tissue. Formazan extraction was performed at room temperature, and the tissues were allowed to stand overnight. Subsequently, 200- $\mu$ L extracts were transferred to a 96-well plate. The optical density was measured at 570 nm and 650 nm as a reference absorbance, with isopropanol as a blank.

The tissue viability was calculated as a percentage relative to the viability of the negative controls. The median of three values from identically treated tissues was used to classify a chemical according to the prediction model.

The amount of IL-1 $\alpha$  released in the conditioned medium after 42 hours was determined using an IL-1 $\alpha$  ELISA kit (Invitrogen, CA, USA), following the manufacturer's detailed instructions.

### 7-2. Prediction model of skin irritation

In this study, the prediction model of skin irritation potential with LabCyte EPI-MODEL was set to refer to the conditions for EPISKIN described in the ECVAM Performance Standards. This predic-

tion model is described in Table 3. In the event that the three independent results within an individual batch were not consistent, the result that occurred twice was used.

Acceptance criteria

- 1) OD<sub>NC</sub> of the negative control is greater than 0.7.
- 2) The viability of the positive control is less than 40%.

Table 3. Positive Criteria.

Tissue Viability (primary)	IL-1 $\alpha$ ELISA (secondary)	Classification
Mean tissue viability $\leq$ 50%	Mean IL-1 $\alpha$ release $\geq$ 120 pg/tissue	Irritant
Mean tissue viability > 50%		
Mean tissue viability > 50%	Mean IL-1 $\alpha$ release < 120 pg/tissue	Non-irritant

**7-3. Difference between LabCyte EPI-MODEL 24 protocol and EPISKIN protocol**

The differences between the LabCyte EPI-MODEL 24 protocol and EPISKIN protocol are summarized in Table 3. Although the amount of medium (Table 4(A)), amount of test chemicals (Table 4(B)), and threshold of IL-1 $\alpha$  content (Table 4(C)) for the LabCyte EPI-MODEL 24 protocol are different from the EPISKIN protocol, their conditions meet the descriptions of the Performance Standards.

Table 4. Differences between the LabCyte EPI-MODEL 24 protocol and EPISKIN protocol.

(A) Amount of medium.

	LabCyte EPI-MODEL 24 SOP	EPISKIN SOP	Reason
Pre-incubation	0.5 mL	2 mL	LabCyte EPI-MODEL 24 cultures are performed in 24-well culture plates. A medium volume of 0.5 mL to 1 mL is appropriate to add to the 24-well culture plate. A medium volume of 1 mL is necessary for a 42-hour culture.
Post-incubation	1 mL	2 mL	
MTT assay	0.5 mL	2 mL	

These conditions meet the descriptions of the Performance Standards.

(B) Amount of test chemicals.

Test chemical	LabCyte EPI-MODEL 24 SOP	EPISKIN SOP	Reason
Liquid	25 $\mu$ L (75 $\mu$ L/cm <sup>2</sup> )	10 $\mu$ L (25 $\mu$ L/cm <sup>2</sup> )	The lowest amount of the test chemical that spread uniformly was applied to the test model.
Solid	25 mg+25 $\mu$ L DW (75 $\mu$ L/cm <sup>2</sup> )	10 mg+10 $\mu$ L DW (25 $\mu$ L/cm <sup>2</sup> )	

These conditions meet the descriptions of the Performance Standards.

(C) Amount of test chemicals.

LabCyte EPI-MODEL 24 SOP	EPISKIN SOP	Performance Standards (EPISKIN)
IL-1 $\alpha$ content $\geq$ 120 pg/tissue (IL-1 $\alpha$ content $\geq$ 120 pg/mL)	IL-1 $\alpha$ content $\geq$ 100 pg/tissue (IL-1 $\alpha$ content $\geq$ 50 pg/mL)	IL-1 $\alpha$ content $\geq$ 120 pg/tissue (IL-1 $\alpha$ $\geq$ 60 pg/mL)

The threshold of IL-1 $\alpha$  released in LabCyte EPI-MODEL was set based on the conditions for EPISKIN described in the Performance Standards.

#### 7-4. Data collection, handling, and analysis

The independent biostatisticians for the study collected and organised the data using specific data collection software (Datasheet4.0:20080910.xls in 2<sup>nd</sup> phase study and Datasheet5.0:20090430.xls in 3<sup>rd</sup> phase study). They will work in close collaboration with the biostatisticians, (Takashi Omori, Etsuyoshi Miyaoka, and Kenya Ishiyama). After decoding the data, they will perform statistical analyses. The data management procedures and statistical tools applied will be approved by the VMT.

#### 7-5. Quality assurance, GLP

##### Laboratories

All participating laboratories worked in the spirit of OECD GLP-like principles.

##### QA aspects

Takashi Omori, Kenya Ishiyama and Hajime Kojima assured the quality of all the data and records.

### 8. Results

#### 8-1 1<sup>st</sup> Phase

##### 8-1-1 Negative control

In 1<sup>st</sup> phase data, Table 5 shows the absorbance values for the negative control. All data for the negative control met the acceptance criteria.

Table 5. Absorbance of negative control by 1<sup>st</sup> phase study.

Lab.	Exp.			Mean	SD
	1	2	3		
a	1.073	0.928	1.007	1.003	0.073
b	0.93	1.245	1.042	1.072	0.16
c	0.96	0.869	0.761	0.863	0.1
d	0.987	0.928	0.939	0.951	0.031
e	0.84	0.884	0.973	0.899	0.068
f	1.049	0.934	0.968	0.984	0.059
g	1.147	1.159	1.074	1.127	0.046

### 8-1-2 Positive control and test chemicals

Table 6 shows the testing chemicals did not show any great score when the scores on tests were repeated in each laboratory. Furthermore, there was no significant inter-laboratory variation. These experiments suggested the feasibility of the LabCyte EPI-MODEL24 through the experiment. All laboratories were judged to participate at the Phase II by the validation management team.

Table 6. Viability of the positive control and three coded chemicals by 1<sup>st</sup> phase study

Chem.	Lab.	1	2	3	Mean	SD
		Viability	Viability	Viability		
PC	a	6.35	27.55	15.67	16.52	10.63
	b	3.94	3.51	3.97	3.81	0.26
	c	5.45	4.81	3.49	4.58	1
	d	11.74	7.22	14.08	11.02	3.49
	e	31.6	9.76	38.61	26.66	15.05
	f	3.1	2.89	2.93	2.97	0.11
	g	4.46	7.17	2.62	4.75	2.29
P01	a	62.67	39.12	46.61	49.46	12.03
Ethanol	b	41.08	50.86	86.58	59.51	23.95
	c	68.13	34.13	67.31	56.53	19.4
	d	68.57	40.52	33.03	47.37	18.73
	e	54.19	72.08	60.55	62.27	9.07
	f		64.16	47.98	56.07	11.44
	g	4.68	5.23	6.67	5.53	1.03
	P02	a	103.63	104.17	98.48	102.09
Glycerol	b	85.5	100.58	67.97	84.68	16.32
	c	101.24	99.41	104.84	101.83	2.76
	d	103.3	101.35	89.73	98.13	7.34
	e	101.75	98.06	99.04	99.62	1.91
	f		97.23	96	96.62	0.87
	g	94	98.16	103.6	98.59	4.82
	P03	a	109.13	90.73	97.78	99.22
naphtalen acetic acid	b	93.96	103.91	103.96	100.61	5.76
	c	103.66	102.11	117.3	107.69	8.36
	d	102.28	98.15	94.56	98.33	3.86
	e	107.11	104.39	97.36	102.95	5.03
	f		101.34	102.07	101.7	0.52
	g	92.2	101.04	105.52	99.59	6.78

### 8-2. 2<sup>nd</sup> phase & 3<sup>rd</sup> phase

#### 8-2-1. Comments at the Datasheet

All tests were sufficient with acceptance criteria. There were a few comments from each laboratory in Tables 7 -9. By an application of Potassium hydroxide (5%aq) (B175, D213 and F232), the model's layers were desquamated. By an application of cinnamaldehyde (D216 and G256), the cups were discoloured and crystallized.



Table 7. Comments on the datasheets (Viability) by 2<sup>nd</sup> phase

Lab ID	Exp.No.	Lot	Date	Comments
a	Main-2	LCE24-081013-B	2008/10/20	This test was recorded as the
a	Main-3	LCE24-081117-B	2008/11/1	This test was recorded as the
a	Main-4	LCE24-081117-B	2008/11/22	This test was recorded as the
b	Main-1	LCE24-081013-B	2008/10/20	
b	Main-2	LCE24-081027-B	2008.11.04	
b	Main-3	LCE24-081117-B	2008/11/25	
c	1	LCE24-080929-B	2008.10.6	
c	2	LCE24-081020-B	2008/10/27	
c	3	LCE24-081027-B	2008.11.3	
d	81021	LCE24-081020-B	2008/10/27	
d	81028	LCE24-081027-B	2008/11/4	
d	81118	LCE24-081117-B	2008/11/25	
e	Main-1	LCE24-081006-B	2008/10/14	
e	Main-2	LCE24-081013-B	2008/10/20	
e	Main-3	LCE24-081020-B	2008/10/27	
f	LAB-08VAL	LCE24-080929-B	2008/10/6	
f	Maruishi	LCE24-081013-B	2008/10/20	
f	LAB-08VAL	LCE24-081103-B	2008/11/10	
g	Main-1	LCE24-080929-B	2008.10.06	By an application of G49,G53,G55, the model's cap was
g	Main-2	LCE24-081013-B	2008.10.20	By an application of G49,G53,G55, the model's cap was
g	Main-3	LCE24-081027-B	2008.11.03	By an application of G49,G53,G55, the model's cap was

Table 8. Comments on the datasheets (ELISA) by 2<sup>nd</sup> phase

Lab ID	Exp.No.	Lot	Date	Comments
a	Main-2	LCE24-081013-B	2008/10/20	This test was recorded as the
a	Main-3	LEC24-081117-B	2008/11/1	This test was recorded as the
a	Main-4	LCE24-081117-B	2008/11/22	This test was recorded as the
b	Main-1	LCE24-081013-B	2008/12/12	
b	Main-2	LCE24-081027-B	2008/12/12	
b	Main-3	LCE24-081117-B	2008.12.26	
c	1	LCE24-080929-B	2008/10/7	
c	2	LCE24-081020-B	2008/10/30	
c	3	LCE24-081027-B	2008.11.3	
d	81021	LCE24-081020-B	2008/11/11	
d	81028	LCE24-081027-B	2008/11/26	
d	81118	LCE24-081117-B	2009/1/7	
e	Main-1	LCE24-081006-B	2008/12/2	
e	Main-2	LCE24-081013-B	2008/12/2	
e	Main-3	LCE24-081020-B	2008/12/19	
f	Maruishi	LCE24-081013-B	2008/11/25	
f	Maruishi	LCE24-081013-B	2008/11/27	
f	LAB-08VAL	LCE24-081103-B	2008/12/25	
g	Main-1	LCE24-080929-B	2008.10.09	
g	Main-2	LCE24-081013-B	2008.10.22	
g	Main-3	LCE24-081027-B	2008.11.05	

Table 9. Comments on the datasheets (Viability) by 3<sup>rd</sup> phase study

Lab ID	Exp.No.	Lot	Date	Comments
a	No.1	LCE24-090420-A	2009/4/27	
a	No.2	LEC24-090511-A	2009/5/18	
a	No.3	LEC24-090518-A	2009/5/25	
b	20090421-1	LCE24-090420-A	2009/4/27	By an application of B175, the model's layers were desquamated.
b	20090421-2	LEC24-090511-A	2009/5/20	By an application of B175, the model's layers were desquamated.
b	20090421-3	LEC24-090518-A	2009/5/25	By an application of B175, the model's layers were desquamated.
c	1	LCE24-090420-A	2009/4/27	
c	2	LEC24-090511-A	2009/5/18	
c	3	LEC24-090518-A	2009/5/25	
d	90512	LEC24-090511-A	2009/5/18	By an application of D213, the model's layers were desquamated. By an application of D216, white crystallizations in the cup were detected.
d	90519	LEC24-090518-A	2009/5/25	By an application of D213, the model's layers were desquamated. By an application of D216, white crystallizations in the cup were detected.
d	90526	LEC24-090525-A	2009/6/1	By an application of D213, the model's layers were desquamated. By an application of D216, white crystallizations in the cup were detected.
f	LAB-09VAL	LCE24-090420-A	2009/4/27	By an application of F232, the model's layers were desquamated.
f	LAB-09VAL	LEC24-090511-A	2009/5/18	
f	LAB-09VAL	LEC24-090518-A	2009/5/25	By an application of F232, the model's layers were desquamated.
g	①	LCE24-090420-A	2009/4/27	By an application of G256, the model's caps were discolored.
g	②	LCE24-090427-A	2009/5/4	By an application of G256, the model's caps were discolored.
g	③	LEC24-090511-A	2009/5/18	By an application of G256, the model's caps were discolored.

**8-2-2. Negative control**

In Table 10 and Fig.2, absorbances of negative control are shown. All data of negative control were sufficient with acceptance criteria excluding Lab a, test1. The mean OD of lab a, test 1 is 0.59 (0.61, 0.58, 0.57). We were not accepted at this result, and accepted the results of test 2-4 re-tested at Lab a.

Table 10 Absorbance of negative control

Study	Run	Lab.					
		a	b	c	d	f	g
2	1	0.75 (0.02)	0.93 (0.01)	0.91 (0.01)	0.82 (0.02)	0.84 (0.01)	1.13 (0.01)
	2	0.86 (0.02)	0.85 (0.04)	1.01 (0.02)	0.90 (0.04)	0.79 (0.02)	1.18 (0.02)
	3	0.82 (0.04)	0.84 (0.03)	0.93 (0.02)	0.96 (0.03)	0.83 (0.00)	1.05 (0.05)
3	1	0.90 (0.02)	0.96 (0.02)	1.04 (0.02)	1.11 (0.05)	0.90 (0.02)	0.91 (0.04)
	2	0.72 (0.02)	1.01 (0.02)	1.06 (0.01)	1.11 (0.04)	0.94 (0.02)	1.08 (0.01)
	3	0.80 (0.02)	0.97 (0.04)	1.01 (0.02)	1.13 (0.03)	0.92 (0.03)	0.88 (0.03)
Mean		0.81	0.93	0.99	1.01	0.87	1.04
Median		0.81	0.94	1.01	1.03	0.87	1.06
Min		0.72	0.84	0.91	0.82	0.79	0.88
Max		0.9	1.01	1.06	1.13	0.94	1.18
SD		0.07	0.07	0.06	0.13	0.06	0.12
Range		0.17	0.17	0.15	0.31	0.15	0.3

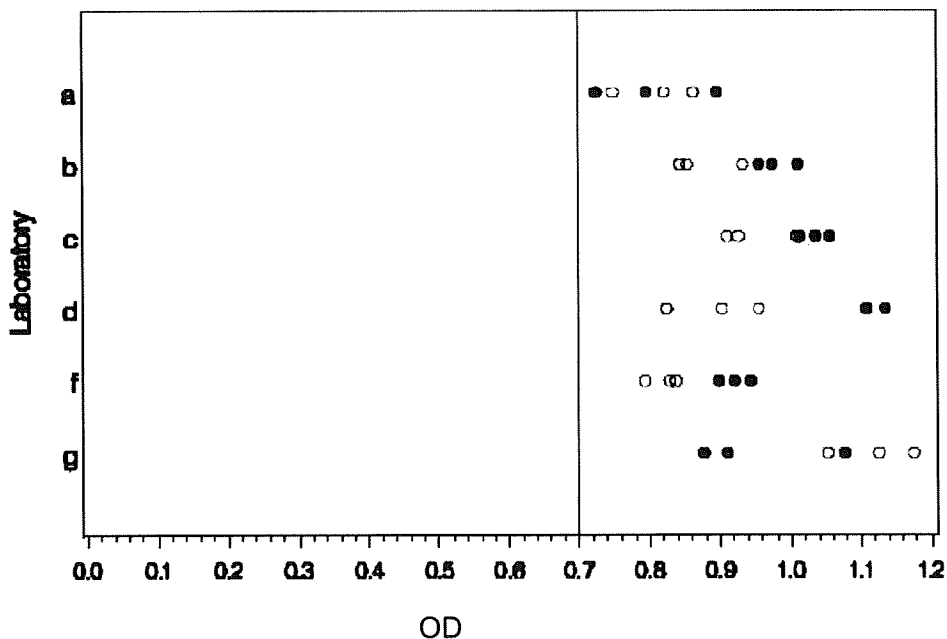


Fig.2 Distribution of Absorbance on negative control

**8-2-3. Positive control**

Table 11 and Fig.3 show three independent viabilities and statistical analysis of positive control at each laboratory. All data were sufficient with acceptance criteria of positive control.

Table 11. Viability of positive control

Study	Run	Lab.					
		a	b	c	d	f	g
2	1	5.9 (1.3)	5.2 (2.3)	4.1 (0.5)	5.7 (2.3)	3.5 (0.4)	3.1 (0.2)
	2	8.8 (4.8)	12.3 (6.9)	5.4 (3.0)	2.6 (0.3)	2.9 (0.3)	10.7 (5.3)
	3	2.5 (0.4)	7.8 (2.4)	3.8 (0.0)	3.3 (0.3)	3.2 (0.3)	4.2 (1.3)
3	1	6.4 (1.8)	9.3 (6.8)	8.2 (3.4)	3.5 (0.9)	8.5 (1.9)	11.7 (2.5)
	2	2.2 (0.4)	2.2 (0.1)	7.3 (2.2)	2.5 (0.3)	4.1 (1.3)	2.5 (0.1)
	3	1.8 (0.2)	1.6 (0.3)	2.4 (0.2)	2.1 (0.4)	2.7 (0.0)	3.3 (0.3)
Mean		4.6	6.4	5.2	3.3	4.1	5.9
Median		4.2	6.5	4.7	2.9	3.3	3.7
Min		1.8	1.6	2.4	2.1	2.7	2.5
Max		8.8	12.3	8.2	5.7	8.5	11.7
SD		2.9	4.2	2.2	1.3	2.2	4.1
Range		7	10.7	5.7	3.6	5.8	9.2

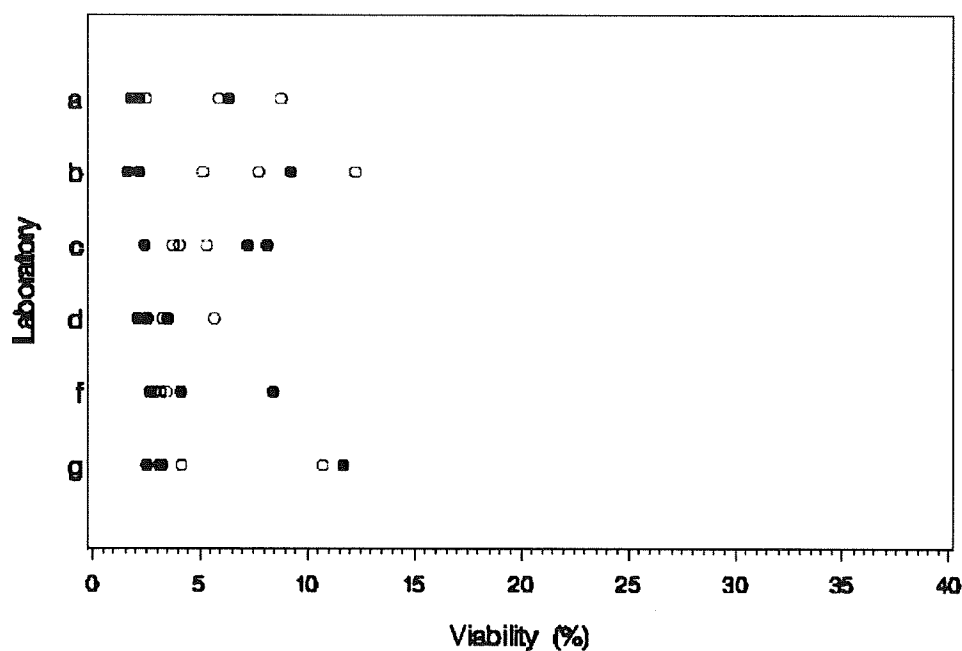


Fig.3 Distribution of viability on positive control

#### 8-2-4. Skin irritation test by cell viability

The results of the skin irritation test with LabCyte EPI-MODEL 24 when it was only evaluated cell viabilities as indicator are shown in Table 12 in 2<sup>nd</sup> phase study and Table 14 in 3<sup>rd</sup> phase study. Summary statistical analysis of viability each chemical are shown in Table 13 and Fig.4 in 2<sup>nd</sup> phase study and Table 15 and Fig.5 in 3<sup>rd</sup> phase study.

Invalid data obtained only Lab a, run 1. This lab performed at retesting. Therefore, the data of lab a were accepted among run 2-4.

Table 12. Viability of chemicals at each laboratory by 2<sup>nd</sup> phase study.

Chem.	Vivo	Score	Exp.	Lab.						
				a	b	c	d	e	f	g
01	no	0	1	31.0	47.1	10.6	14.3	38.1	14.3	10.6
			2	11.2	10.4	20.3	9.1	25.2	11.2	10.6
			3	11.6	16.1	12.4	9.6	32.3	10.4	14.0
02	no	0	1	79.8	66.9	88.1	102.3	101.8	75.3	96.0
			2	76.5	61.7	89.7	89.8	76.4	67.2	94.8
			3	65.2	88.7	85.8	67.6	85.8	75.7	103.3
03	no	0	1	109.1	93.3	94.6	105.1	129.6	94.2	100.5
			2	103.9	99.8	93.1	112.8	106.6	97.9	93.4
			3	100.9	102.3	95.7	101.4	103.9	92.5	111.1
04	no	0	1	106.3	94.4	97.1	106.1	127.1	100.1	104.8
			2	95.2	100.2	99.9	100.9	113.6	92.8	103.3
			3	96.5	98.6	97.8	98.4	105.2	92.7	109.8
05	no	0.3	1	78.5	61.7	91.4	79.4	103.0	71.9	96.8
			2	78.5	71.9	95.2	70.5	90.3	39.3	89.9
			3	74.1	84.5	89.2	66.1	89.6	55.1	88.4
06	no	0.3	1	92.5	77.9	81.0	91.3	97.0	87.8	87.2
			2	79.4	83.5	79.1	102.4	81.5	94.4	81.2
			3	82.4	80.5	83.6	82.7	90.7	81.1	54.1
07	no	1	1	24.1	10.8	20.8	21.7	17.5	15.8	31.5
			2	12.6	12.6	16.2	13.8	22.2	31.1	22.5
			3	17.8	13.2	15.2	19.8	21.3	15.6	19.9
08	no	1	1	111.9	86.7	75.3	109.4	114.9	89.7	101.1
			2	90.2	100.6	82.3	107.5	100.9	97.8	100.9
			3	95.3	104.8	77.2	103.0	100.9	96.5	109.0
09	no	1.7	1	112.8	96.7	106.6	105.0	115.8	98.8	102.3
			2	97.1	110.1	96.8	103.4	108.6	86.5	103.4
			3	101.1	109.5	93.5	98.1	103.9	97.7	112.1
10	no	1.7	1	115.9	115.4	107.5	114.3	132.0	104.0	107.9
			2	104.1	110.1	103.6	108.2	117.0	101.2	108.4
			3	86.5	111.3	103.7	105.5	107.5	101.2	113.1

Table 12. continued

Chem.	Vivo	Score	Exp.	Lab.						
				a	b	c	d	e	f	g
11	no	2	1	113.7	105.0	101.0	102.4	123.1	103.1	102.8
			2	98.1	106.6	94.6	105.8	110.4	98.0	100.5
			3	112.6	103.7	94.1	102.7	105.5	94.6	109.0
12	no	2	1	28.2	24.6	24.9	54.3	55.6	27.2	87.7
			2	18.4	24.6	44.8	76.2	57.8	65.2	98.0
			3	15.3	15.9	28.1	27.4	57.2	66.0	112.6
14	Category 2	2.3	1	11.1	12.1	14.7	10.7	14.2	13.1	13.5
			2	6.6	8.3	9.5	11.7	12.0	16.7	12.0
			3	6.8	8.8	9.1	10.2	10.4	17.0	10.6
15	Category 2	2.3	1	11.1	9.3	13.1	8.0	11.0	8.6	9.2
			2	7.1	10.2	19.3	8.6	11.3	5.9	24.7
			3	8.2	9.9	8.1	9.2	8.7	7.1	9.2
16	Category 2	2.7	1	67.9	92.0	51.5	18.1	98.2	59.6	64.9
			2	32.2	54.1	86.3	79.2	90.6	50.4	79.6
			3	59.8	98.3	81.7	37.7	78.7	67.5	86.5
17	Category 2	2.7	1	6.1	4.5	5.3	6.6	8.9	6.9	6.2
			2	4.8	4.7	6.0	5.3	6.3	5.5	5.3
			3	5.6	5.7	5.9	3.9	5.4	4.5	5.3
18	Category 2	3	1	82.1	46.5	91.2	83.7	98.9	69.2	92.4
			2	78.3	50.6	87.3	69.9	87.2	80.6	85.9
			3	25.3	100.0	87.5	59.0	69.1	71.9	94.4
19	Category 2	3	1	15.0	74.6	10.0	30.4	83.1	40.1	35.8
			2	19.9	10.9	22.4	28.3	26.1	87.0	44.7
			3	51.1	32.0	35.0	18.2	69.4	71.8	38.7
20	Category 2	4	1	31.1	24.8	10.4	9.6	10.7	8.1	8.8
			2	9.3	8.0	7.6	16.9	8.2	7.8	6.7
			3	29.5	9.3	7.6	30.9	6.2	8.2	8.6

Table 13. Summary of the statistical analysis of the viability for each chemical by 2<sup>nd</sup> phase study.

Chem.	Stat.	Lab.						
		a	b	c	d	e	f	g
01	Mean	17.9	24.5	14.4	11.0	31.9	12.0	11.7
	Median	11.6	16.1	12.4	9.6	32.3	11.2	10.6
	Min	11.2	10.4	10.6	9.1	25.2	10.4	10.6
	Max	31.0	47.1	20.3	14.3	38.1	14.3	14.0
02	Mean	73.8	72.4	87.8	86.6	88.0	72.7	98.0
	Median	76.5	66.9	88.1	89.8	85.8	75.3	96.0
	Min	65.2	61.7	85.8	67.6	76.4	67.2	94.8
	Max	79.8	88.7	89.7	102.3	101.8	75.7	103.3
03	Mean	104.7	98.5	94.5	106.4	113.3	94.8	101.7
	Median	103.9	99.8	94.6	105.1	106.6	94.2	100.5
	Min	100.9	93.3	93.1	101.4	103.9	92.5	93.4
	Max	109.1	102.3	95.7	112.8	129.6	97.9	111.1
04	Mean	99.3	97.8	98.2	101.8	115.3	95.2	105.9
	Median	96.5	98.6	97.8	100.9	113.6	92.8	104.8
	Min	95.2	94.4	97.1	98.4	105.2	92.7	103.3
	Max	106.3	100.2	99.9	106.1	127.1	100.1	109.8
05	Mean	77.0	72.7	91.9	72.0	94.3	55.4	91.7
	Median	78.5	71.9	91.4	70.5	90.3	55.1	89.9
	Min	74.1	61.7	89.2	66.1	89.6	39.3	88.4
	Max	78.5	84.5	95.2	79.4	103.0	71.9	96.8
06	Mean	84.8	80.7	81.2	92.1	89.7	87.8	74.2
	Median	82.4	80.5	81.0	91.3	90.7	87.8	81.2
	Min	79.4	77.9	79.1	82.7	81.5	81.1	54.1
	Max	92.5	83.5	83.6	102.4	97.0	94.4	87.2
07	Mean	18.2	12.2	17.4	18.4	20.3	20.8	24.6
	Median	17.8	12.6	16.2	19.8	21.3	15.8	22.5
	Min	12.6	10.8	15.2	13.8	17.5	15.6	19.9
	Max	24.1	13.2	20.8	21.7	22.2	31.1	31.5
08	Mean	99.1	97.4	78.3	106.6	105.6	94.7	103.7
	Median	95.3	100.6	77.2	107.5	100.9	96.5	101.1
	Min	90.2	86.7	75.3	103.0	100.9	89.7	100.9
	Max	111.9	104.8	82.3	109.4	114.9	97.8	109.0
09	Mean	103.7	105.4	98.9	102.2	109.4	94.3	105.9
	Median	101.1	109.5	96.8	103.4	108.6	97.7	103.4
	Min	97.1	96.7	93.5	98.1	103.9	86.5	102.3
	Max	112.8	110.1	106.6	105.0	115.8	98.8	112.1
10	Mean	102.1	112.2	104.9	109.3	118.8	102.1	109.8
	Median	104.1	111.3	103.7	108.2	117.0	101.2	108.4
	Min	86.5	110.1	103.6	105.5	107.5	101.2	107.9
	Max	115.9	115.4	107.5	114.3	132.0	104.0	113.1

Table 13. continued.

Chem.	Stat.	Lab.						
		a	b	c	d	e	f	g
11	Mean	108.1	105.1	96.6	103.6	113.0	98.6	104.1
	Median	112.6	105.0	94.6	102.7	110.4	98.0	102.8
	Min	98.1	103.7	94.1	102.4	105.5	94.6	100.5
	Max	113.7	106.6	101.0	105.8	123.1	103.1	109.0
12	Mean	20.7	21.7	32.6	52.6	56.9	52.8	99.5
	Median	18.4	24.6	28.1	54.3	57.2	65.2	98.0
	Min	15.3	15.9	24.9	27.4	55.6	27.2	87.7
	Max	28.2	24.6	44.8	76.2	57.8	66.0	112.6
14	Mean	8.2	9.7	11.1	10.9	12.2	15.6	12.0
	Median	6.8	8.8	9.5	10.7	12.0	16.7	12.0
	Min	6.6	8.3	9.1	10.2	10.4	13.1	10.6
	Max	11.1	12.1	14.7	11.7	14.2	17.0	13.5
15	Mean	8.8	9.8	13.5	8.6	10.3	7.2	14.4
	Median	8.2	9.9	13.1	8.6	11.0	7.1	9.2
	Min	7.1	9.3	8.1	8.0	8.7	5.9	9.2
	Max	11.1	10.2	19.3	9.2	11.3	8.6	24.7
16	Mean	53.3	81.4	73.1	45.0	89.1	59.1	77.0
	Median	59.8	92.0	81.7	37.7	90.6	59.6	79.6
	Min	32.2	54.1	51.5	18.1	78.7	50.4	64.9
	Max	67.9	98.3	86.3	79.2	98.2	67.5	86.5
17	Mean	5.5	4.9	5.8	5.3	6.9	5.6	5.6
	Median	5.6	4.7	5.9	5.3	6.3	5.5	5.3
	Min	4.8	4.5	5.3	3.9	5.4	4.5	5.3
	Max	6.1	5.7	6.0	6.6	8.9	6.9	6.2
18	Mean	61.9	65.7	88.7	70.9	85.1	73.9	90.9
	Median	78.3	50.6	87.5	69.9	87.2	71.9	92.4
	Min	25.3	46.5	87.3	59.0	69.1	69.2	85.9
	Max	82.1	100.0	91.2	83.7	98.9	80.6	94.4
19	Mean	28.7	39.2	22.5	25.6	59.5	66.3	39.8
	Median	19.9	32.0	22.4	28.3	69.4	71.8	44.7
	Min	15.0	10.9	10.0	18.2	26.1	40.1	35.8
	Max	51.1	74.6	35.0	30.4	83.1	87.0	44.7
20	Mean	23.3	14.0	8.6	19.2	8.4	8.0	8.1
	Median	29.5	9.3	7.6	16.9	8.2	8.1	8.6
	Min	9.3	8.0	7.6	9.6	6.2	7.8	6.7
	Max	31.1	24.8	10.4	30.9	10.7	8.2	8.8



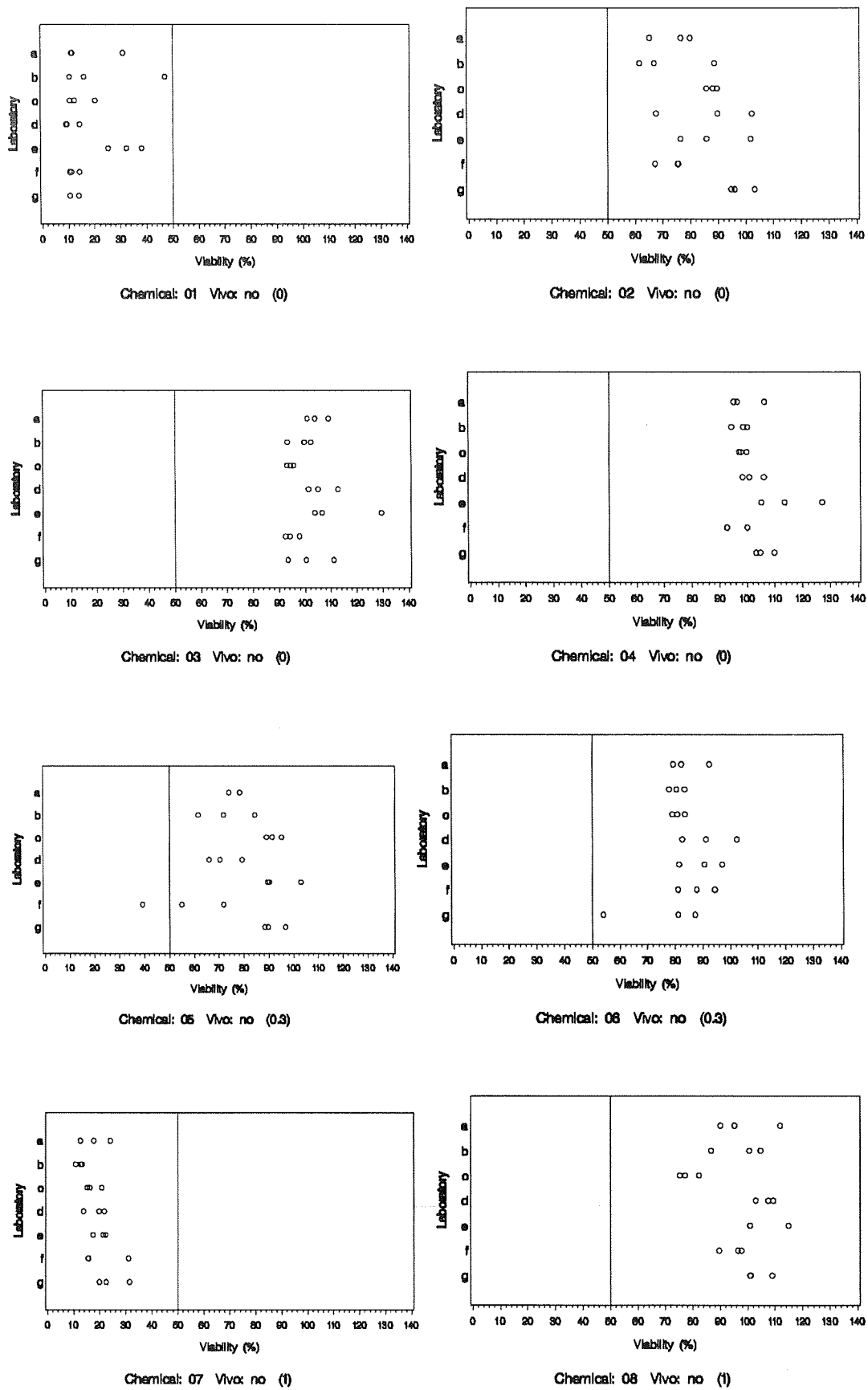
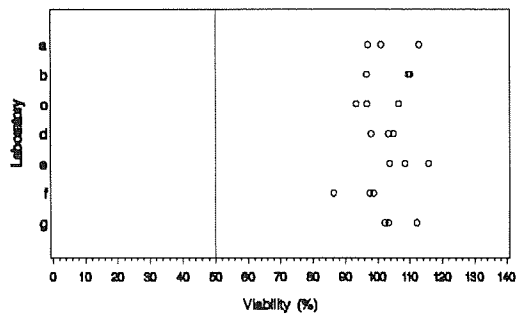
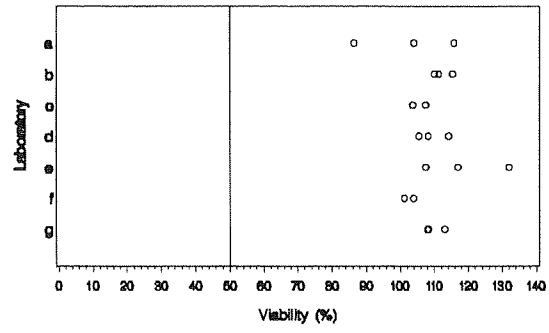


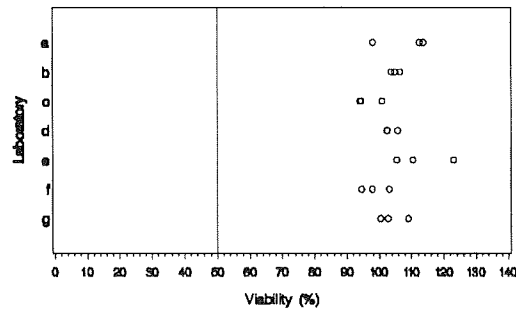
Fig. 4. Distribution of the viability for each chemical.



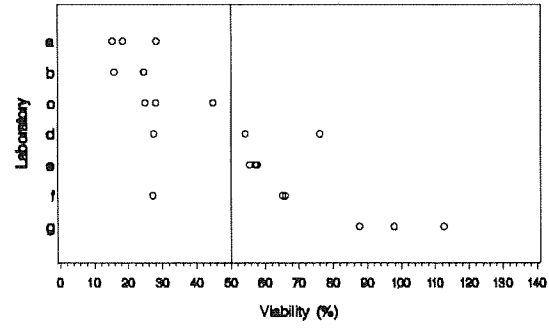
Chemical: 09 Vivo: no (1.7)



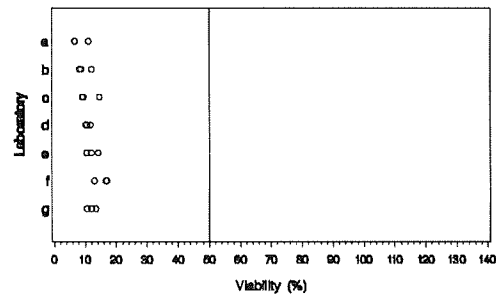
Chemical: 10 Vivo: no (1.7)



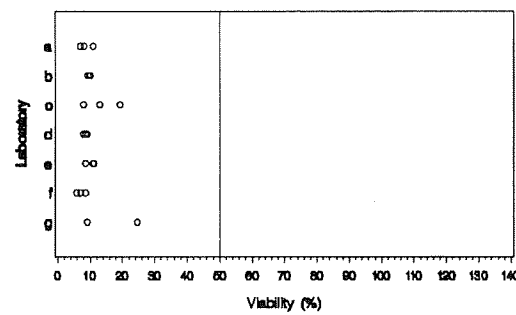
Chemical: 11 Vivo: R38 (2)



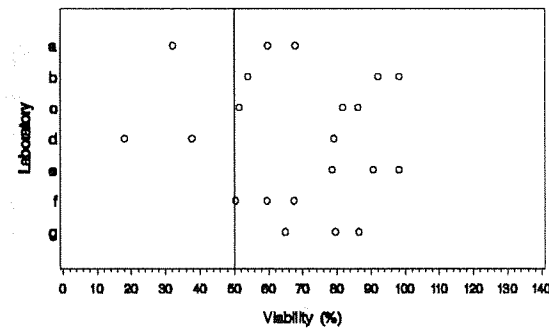
Chemical: 12 Vivo: R38 (2)



Chemical: 14 Vivo: R38 (2.3)



Chemical: 15 Vivo: R38 (2.3)



Chemical: 18 Vivo: R38 (2.7)

Fig. 4. continued

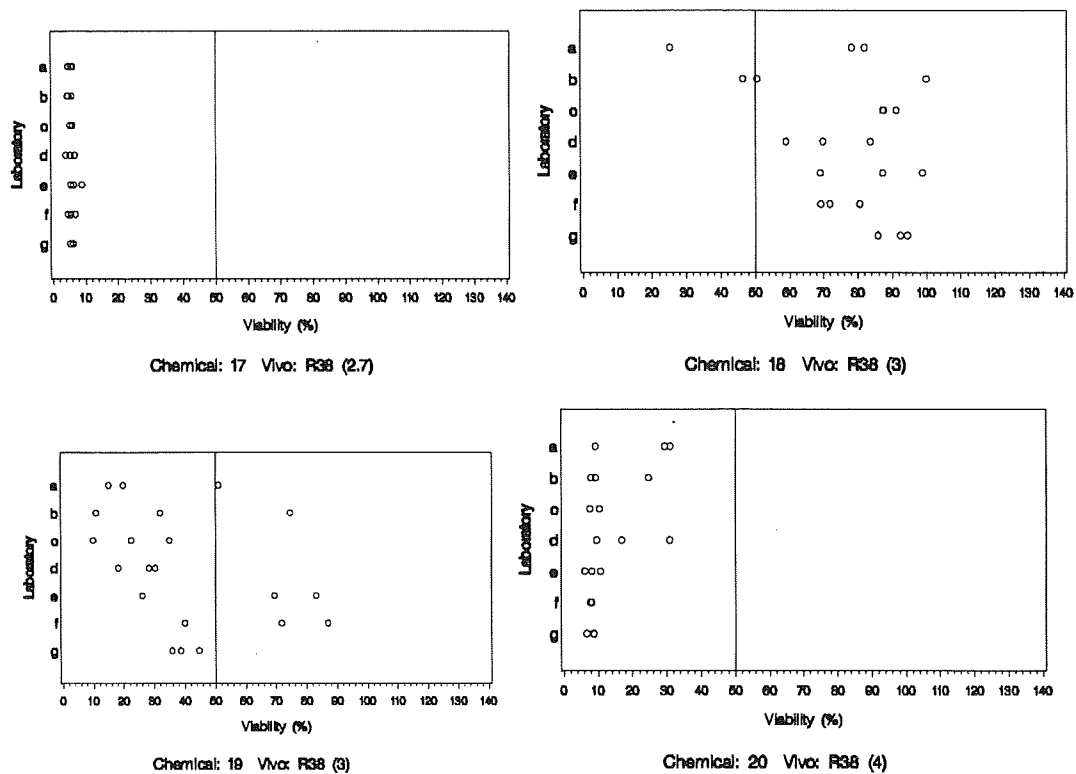


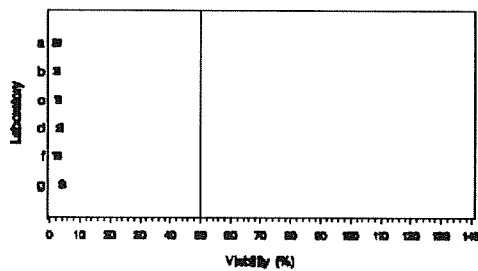
Fig. 4. continued.

Table 14. Viability of chemicals each laboratory by 3<sup>rd</sup> phase study

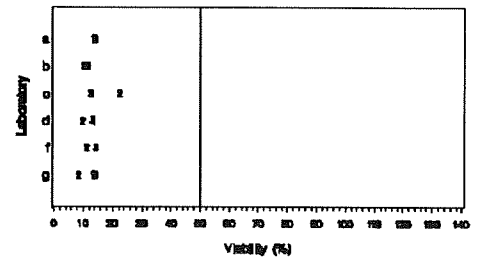
Chem.	Vivo	Score	Exp.	Lab.					
				a	b	c	d	f	g
A	no	2	1	13.3	11.8	13.2	13.8	11.4	13.7
			2	14.2	10.2	22.5	9.9	11.3	8.7
			3	14.0	11.1	12.3	13.2	14.3	14.3
B	Category 2	2.7	1	1.5	2.2	2.5	4.0	1.7	3.9
			2	3.1	2.2	2.9	3.0	2.6	3.7
			3	1.5	2.5	3.0	3.9	3.2	4.7
C	Category 2	3	1	0.7	0.7	0.7	6.9	0.8	1.0
			2	1.3	1.1	1.4	2.0	4.8	0.4
			3	0.5	0.8	1.0	0.8	1.0	0.3
D	Category 2	3.3	1	14.5	24.0	12.7	10.3	13.8	19.3
			2	13.6	16.0	12.5	18.3	8.8	15.2
			3	18.6	15.5	12.6	23.0	19.2	14.1
E	Category 2	3.3	1	3.9	3.4	3.4	8.2	3.2	4.1
			2	4.5	2.7	3.3	3.9	4.2	3.1
			3	1.8	3.5	3.5	3.7	5.0	5.1
F	Category 2	4	1	5.6	7.2	6.5	6.4	5.2	7.2
			2	5.7	6.1	6.8	5.4	7.4	6.8
			3	5.4	4.2	6.5	5.4	5.0	7.6

Table 15 Summary statistical analysis of viability each chemical by 3<sup>rd</sup> phase study

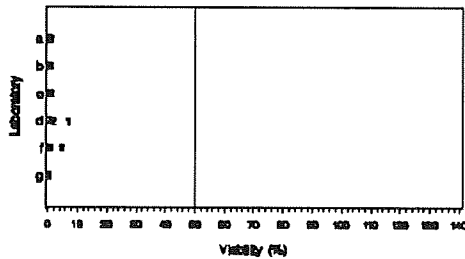
Chem.	Stat.	Lab.					
		a	b	c	d	f	g
A	Mean	13.8	11.0	16.0	12.3	12.3	12.2
	Median	14.0	11.1	13.2	13.2	11.4	13.7
	Min	13.3	10.2	12.3	9.9	11.3	8.7
	Max	14.2	11.8	22.5	13.8	14.3	14.3
B	Mean	2.0	2.3	2.8	3.6	2.5	4.1
	Median	1.5	2.2	2.9	3.9	2.6	3.9
	Min	1.5	2.2	2.5	3.0	1.7	3.7
	Max	3.1	2.5	3.0	4.0	3.2	4.7
C	Mean	0.8	0.8	1.0	3.2	2.2	0.6
	Median	0.7	0.8	1.0	2.0	1.0	0.4
	Min	0.5	0.7	0.7	0.8	0.8	0.3
	Max	1.3	1.1	1.4	6.9	4.8	1.0
D	Mean	15.6	18.5	12.6	17.2	13.9	16.2
	Median	14.5	16.0	12.6	18.3	13.8	15.2
	Min	13.6	15.5	12.5	10.3	8.8	14.1
	Max	18.6	24.0	12.7	23.0	19.2	19.3
E	Mean	3.4	3.2	3.4	5.3	4.2	4.1
	Median	3.9	3.4	3.4	3.9	4.2	4.1
	Min	1.8	2.7	3.3	3.7	3.2	3.4
	Max	4.5	3.5	3.5	8.2	5.0	5.1
F	Mean	5.5	5.8	6.6	5.7	5.9	7.2
	Median	5.6	6.1	6.5	5.4	5.2	7.2
	Min	5.4	4.2	6.5	5.4	5.0	6.8
	Max	5.7	7.2	6.8	6.4	7.4	7.6



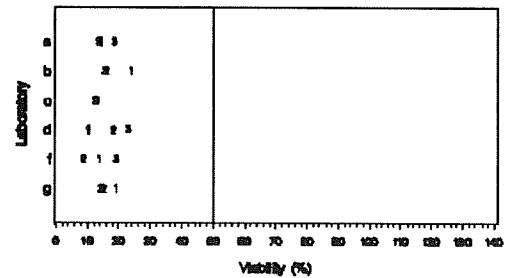
2-chloromethyl-3,5-dimethyl-4-methoxypyridine HCl



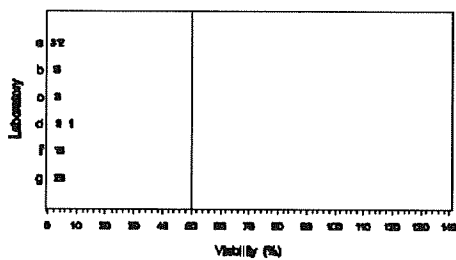
cinnamaldehyde



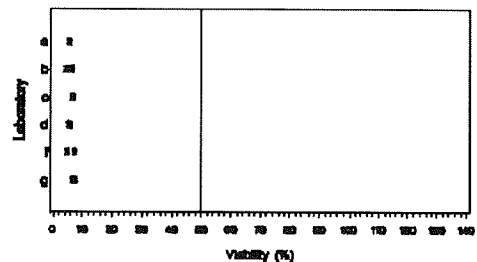
potassium hydroxide (5% aq.)



benzeneethiol, 5-(1,1-dimethylethyl)-2-methyl



1-methyl-3-phenyl-1-piperazine



1,1-bischloroethane

Fig.5 Distribution of viability each chemical