# Annex III: Tabular summary of dose-elicitation studies in sensitised patients

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# Chloroatranol

Chloroatranol (allergen in oak moss absolute: Evernia prunastri) (1)				
Design	blinded, randomised with regard to doses and controlled			
Test subjects	13 patients previously identified as sensitized to chloroatranol and oak moss absolute			
Controls	10 healthy controls			
Substance	Purity: >99%			
Patch test	15 $\mu$ l solution applied in an 8 mm Finn Chamber occlusion 48 h			
-dilution steps	200 ppm to 0.0063 ppm (10 steps)			
-control/vehicle	ethanol			
-definition of threshold	lowest concentration giving a visible skin reaction			
ROAT	volar aspect of forearms			
area	3 x 3 cm <sup>2</sup>			
applications/day	two			
dose	chloroatranol in ethanol: Step 1: 5 ppm Step 2: 25 ppm			
dose/application/cm <sup>2</sup>	step 1: 0.025 µg step2: 0.125 µg			
control substance	ethanol			
definition of positive	erythema in at least 25% and at least one papule			
period	two weeks for each step			
Results				
PT ED10% (95% CI)	0.013 (0.002-0.03) ppm =0.0004 μg/cm <sup>2</sup>			
PT ED50% (95% CI)	0.15 (0.077-0.295) ppm =0.0045 μg/cm <sup>2</sup>			
PT no effect level (observed)	/			
ROAT	Cumulative responses			
Step 1 (5 ppm)	12/13 (92%)			
Step 2 (25 ppm)	13/13 (100%)			
Controls	Negative			
Other information	None relevant			

In a subsequent study chloroatranol and atranol, both ingredients in *Evernia prunastri*, were tested in equimolar concentrations in serial dilution in 10 eczema patients with known sensitization to chloroatranol and oak moss. A positive response was defined as any degree of reaction. Ethanol was included as the control and gave no response. No use tests were done and no control subjects included.

**Results**: All patients reacted to the highest concentrations of the two substances. For both substances there was a significant dose-dependence and the estimated difference in elicitation potency of chloroatranol relative to atranol was 217%. The dose-response curve is seen in figure 1 below (2).



Fig. 1. Observed response rates and fitted parallel logistic dose-response curves for atranol and chloroatranol in equimolar concentrations at patch testing. The response was dichotomized and any reaction other than zero was classified as positive.

# Cinnamal

Cinnamal (3)	
Design	blinded, randomised and controlled
Test subjects	18 patients with a positive patch test to cinnamal and additional 4 with a doubtful response
Controls	20 healthy controls
Substance	Purity: >98%
Patch test	20 mg solution applied in an 8 mm Finn Chamber occlusion 48 h
-dilution steps	2% to 0.01% (7 steps)
-control/vehicle	petrolatum
-definition of threshold	lowest concentration giving a visible skin reaction in a continuous line of responses
ROAT	outer aspect of upper arm
area	5 x 5 cm <sup>2</sup>
applications/day	two with atomizer pump
dose	Step 1: 0.02% Step 2: 0.1% Step 3: 0.8%
dose/application/cm <sup>2</sup>	Not given
control substance	ethanol
definition of positive	The response was classified as positive no matter the degree of reaction.
period	two weeks for each step; total maximum 6 weeks
Results	
PT ED10% (95% CI)	
PT ED50% (95% CI)	0.24% = 96 $\mu$ g/cm <sup>2</sup> (calculated from the data in the paper)
PT no effect level(observed)	0.01 % in pet. = 0.4 $\mu$ g/cm <sup>2</sup>
ROAT	Cumulative responses
Step 1 (0.02%)	0/18
Step 2 (0.1%)	8/18 (44 %)
Step 3 (0.8%)	13/18 (72 %)
Controls	No eczema reactions were seen
Other information	2 patients and 2 controls developed immediate reactions to the cinnamal solution

Cinnamal (4)		
Design	blinded, randomised doses and controlled	
Test subjects	17 patients with a positive patch test to cinnamal (8 patients in part 1 and 9 in part two)	
Controls	20 controls (non-sensitised dermatitis patients)	
Substance	purity: /	
Patch test	15 $\mu$ l solution applied in an 8 mm Finn Chamber occlusion 48 h	
-dilution steps	2 % to 0.00006 % (17 steps)	
-control/vehicle	ethanol	
-definition of threshold	lowest concentration eliciting a + reaction	
ROAT	Axilla	
area	10 x 10 cm2 (estimated)	
applications/day	two with roll on deodorant ( 89-700 mg per application of solution) average cases: 263 mg/application controls: only range given	
dose	Part one: Step 1: 0.032% Step 2: 0.1% Step: 0.32% Part two: Step 1: 0.01% Step 2: 0.032% Step 3: 0.1%	
dose/application/cm2	Part two estimated: step one: 0.26 µg; step two: 0.84 µg; 2.63 µg	
control substance	Deodorant matrix	
definition of positive	eczematous reaction covering at least 25% of test area	
period	Part one: one week with each concentration: maximum three weeks Part two: two weeks with each concentration: maximum six weeks	
Results		
PT ED10% (95% CI)	/	
PT ED50% (95% CI)	/	
PT no effect level(observed)	0.002%	
ROAT	Cumulative responses	
Step 1 (0.01)	2/9 (22%)	
Step 2 (0.032)	6/9 (67%)	
Step 3 (0.1)	8/9 (88%)	
Controls	No reactions were seen	
Other information	Only reactions seen to the cinnamal-containing deodorants at ROAT, difference to matrix axilla ( $p$ <0.001) and all control persons negative ( $p$ <0.001)	

# Hydroxycitronellal

Hydroxycitronellal (5)	
Design	blinded, randomised doses and controlled
Test subjects	7 patients with a positive patch test to hydroxycitronellal
Controls	7 controls (non-sensitised dermatitis patients)
Substance	purity: /
Patch test	15 $\mu$ l solution applied in an 8 mm Finn Chamber occlusion 48 h
-dilution steps	4% to 0.00006% (17 steps)
-control/vehicle	ethanol
-definition of threshold	lowest concentration eliciting + reaction
ROAT	Axilla
area	10 x 10 cm <sup>2</sup> (estimated)
applications/day	two with roll on deodorant (172-591 per application of solution) average cases: 294 mg/application controls: only range given
dose	Step 1: 0.032% Step 2: 0.1% Step: 0.32%
dose/application/cm <sup>2</sup>	Estimated: step 1: 0.94 µg; step 2: 2.94 µg; step 3: 9.40 µg
control substance	Deodorant matrix
definition of positive	eczematous reaction covering at least 25% of test area
period	two weeks with each concentration: maximum six weeks
Results	
PT ED10% (95% CI)	
PT ED50% (95% CI)	/
PT no effect level(observed)	<0.00012 %
ROAT	Cumulative responses
Step 1 (0.032)	4/7 (57%)
Step 2 (0.1)	5/7 (71%)
Step 3 (0.32)	7/7 (100%)
Controls	No reactions were seen
Other information	Reactions were only seen to the hydroxycitronellal-containing deodorant at ROAT, difference to matrix treated axilla ( $p$ <0.001) and all control persons negative ( $p$ <0.001)

Hydroxycitronellal (6)	
Design	double blinded, randomised
Test subjects	13 patients with a positive patch test to hydroxycitronellal
Controls	/
Substance	purity: unknown
Patch test	confirmatory
-dilution steps	
-control/vehicle	
-definition of threshold	
ROAT	finger immersion in fragrance solution in 10% ethanol
area	/
applications/day	Once per day for 10 min
dose	Step 1: 10 ppm Step 2: 250 ppm
dose/application/cm <sup>2</sup>	Not applicable
control substance	10% alcohol
definition of positive	clinical grading scale and laser doppler comparison between active and control
period	two weeks with each concentration: maximum four weeks
Results	
PT ED10% (95% CI)	Not relevant
PT ED50% (95% CI)	Not relevant
PT no effect level(observed)	Not relevant
ROAT	Cumulative responses
Step 1 (10 ppm)	1/13
Step 2 (250 ppm)	5/13
Vehicle control	4/13
Other information	No difference between active substance and control application was found.

Hydroxyisohexyl 3-	-cyclohexenecarboxaldehyde	(HICC)
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Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) (7)				
Design	blinded, randomised and controlled			
Test subjects	18 patients with a positive patch test to HICC			
Controls	7 healthy controls			
Substance	Purity: >99%			
Patch test	15 $\mu$ l solution applied in an 8 mm Finn Chamber occlusion 48 h			
-dilution steps	6% to 0.0006%			
-control/vehicle	ethanol			
-definition of threshold	lowest concentration giving a visible skin reaction in a continuous line of reactions			
ROAT	volar aspect of lower arm			
area	3 x 3 cm <sup>2</sup>			
applications/day	two with droplet bottle (theoretical:30 mg per application of solution)			
dose	Step 1: 0.5% Step 2: 3%			
µg/application/cm <sup>2</sup>	Step 1: 15.3 (3.4-22.2) Step 2: 126.2 (40.5-226.2)			
control substance	ethanol			
definition of positive	erythema in at least 25% and at least one papule			
period	two weeks for each step; total maximum 4 weeks			
Results				
PT ED10% (95% CI)	0.9 μg/cm <sup>2</sup> 29 (7-69) ppm			
PT ED50% (95% CI)	20 μg/cm <sup>2</sup> 662 (350-1250)ppm			
PT no effect level (observed)	/			
ROAT	Cumulative responses			
Step 1 (0.5%)	11/18 (61%)			
Step 2 (3%)	16/18 (89%)			
Controls	No reactions were seen			
Other information	Difference between test and control group statistically significant			

Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) (8)			
Design	blinded, randomised and controlled		
Test subjects	15 patients with a positive patch test to HICC		
Controls	10 healthy controls		
Substance	Purity: > 98.8%		
Patch test	15 $\mu l$ solution applied in an 8 mm Finn Chamber occlusion 48 h		
-dilution steps	6% to 0.0006% (5 steps)		
-control/vehicle	ethanol		
-definition of threshold	lowest concentration giving a visible skin reaction in a continuous line of reactions		
ROAT	Axilla		
area	76 cm <sup>2</sup> (template)		
applications/day	two with roll on deodorant		
dose	Step 1: 200 ppm Step 2: 600 ppm Step 3: 1800 ppm		
dose/application/cm <sup>2</sup>	median 0.79 µg HICC		
control substance	deodorant matrix		
definition of positive	spotty erythema involving at least 25% of the exposed area and infiltration represented by at least one papule.		
period	two weeks for each step; total maximum 6 weeks		
Results			
PT ED10% (95% CI)	0.75 μg/cm <sup>2</sup> 25 ppm (0.69-120)		
PT ED50% (95% CI)	18.3 μg/cm <sup>2</sup> 610 ppm (120-2800)		
PT no effect level (observed)	< 0.0006%		
ROAT	Cumulative responses		
Step 1 (200 ppm)	9/14* (64%)		
Step 2 (600 ppm)	12/14* (86%)		
Step 3 (1800 ppm)	14/14* (100%)		
Controls	No reactions were seen		
Other information	*14 patients completed the use test study Difference between HICC deodorant and matrix deodorant in cases ( $p$ =0.0001).Difference between controls and patients ( $p$ =0.004).		

Hydroxyisohexyl 3-cyclohexenecarboxaldehyde (HICC) (9)			
Design	blinded, randomised and controlled		
Test subjects	17 patients with a positive patch test to HICC		
Controls	15 healthy controls		
Substance	IFF lot SM/8059062		
Patch test	15 µl solution applied in an 8 mm Finn Chamber occlusion 48 h		
-dilution steps	1500 to 0.0022 µg/cm <sup>2</sup> HICC (19 steps)		
-control/vehicle	ethanol		
-definition of threshold	lowest concentration giving a visible skin reaction in a continuous line of reactions to higher concentrations		
ROAT	volar aspect of forearms		
area	3 x 3 cm (5 areas)		
applications/day	two with micropipette (20 $\mu$ l per application)		
dose	Simultaneous application to 5 areas, four doses each and vehicle		
µg /application/cm <sup>2</sup>	Dose 1:0.0357 Dose 2: 0.357 Dose 3: 3.57 Dose 4: 35.7		
control substance	ethanol		
definition of positive	at least 5 points on a clinical scale, corresponding to erythema in 25% of test area and at least 1 papule		
period	Three weeks. All concentrations applied simultaneously (randomised)		
Results			
PT ED10% (95% CI)	0.662 μg/ cm <sup>2</sup> (0.052-2.35)		
PT ED50% (95% CI)	11.1 μg/ cm <sup>2</sup> (3.41- 33.1)		
PT no effect level(observed)	<0.0022 µg/ cm <sup>2</sup>		
ROAT	Cumulative responses		
Dose 1 (0.0357)	0/16*		
Dose 2 (0.357)	3/16 (19%)		
Dose 3 (3.57)	12/16 (75%)		
Dose 4 (35.7)	15/16 (94%)		
Controls	No reactions were seen		
Other information	*16 patients completed the use test study The evaporation rate of HICC was calculated to 72% over a 24-h period. ED10% ROAT: 0.064 $\mu$ g/cm <sup>2</sup> (more info see below)		

Table 2 The dose per application and accumulated dose after 1, 2 and 3 weeks in the ROAT

ROAT, dose per application (µg HICC cm <sup>-2</sup> )	Number of applications after 1 week	Total accumulated dose after 1 week (µg HICC cm <sup>-2</sup> )	Number of applications after 2 weeks	Total accumulated dose after 2 weeks (µg HICC cm <sup>-2</sup> )	Number of applications after 3 weeks	Total accumulated dose after 3 week (µg HICC cm <sup>-2</sup> )
35.7	14	500	28	1000	42	1500
3.57	14	50	28	100	42	150
0.357	14	5	28	10	42	15
0.0357	14	0.5	2.8	1	42	1.5



Fig 3. The fitted dose–response curve for the patch test  $\left(n=16\right)$  and the 1-week, the 2-week 3-week accumulated ROAT doses.

#### Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC)

In a study be the German Contact Dermatitis Group, 64 persons previously diagnosed with HICC contact allergy were exposed to increasing doses of HICC in 2 different formulations, a hydrophilic cream and an ethanol solution, to mimic everyday exposures, following a standardised ROAT protocol (10). The concentration of HICC tolerated by 90% of the sensitised was estimated as 1.2  $\mu$ g/cm<sup>2</sup> for perfume and 4.9  $\mu$ g/cm<sup>2</sup> for cream. The dose-response curve is shown in Fig. 4.3 – 1 below.

Hydroxyisohexyl 3-cyclohex	cene carboxaldehyde (HICC) (10)				
Design	randomised and vehicle controlled				
Test subjects	67 patients with a previous positive patch test to HICC				
Controls	None				
Substance	Provided by International Flavor & Fragrances Inc, Hilversum, NL				
Patch test					
-dilution steps	2.5% and 5%				
-control/vehicle	petrolatum				
-definition of threshold	lowest concentration giving a positive skin reaction in a continuous line to next higher concentration.				
ROAT	Volar forearms (both sides)				
area	3 x 3 cm (4 areas: one test and one control each for alcoholic solution and cream, respectively)				
applications/day	two				
dose	$ \begin{array}{cccccc} 2.8 \ \mu g/cm^2 \ \text{in cream} & 0.2 \ \mu g/cm^2 \ \text{in ethanol} \\ 5.6 \ \mu g/cm^2 \ \text{in cream} & 0.4 \ \mu g/cm^2 \ \text{in ethanol} \\ 55.6 \ \mu g/cm^2 \ \text{in cream} & 4.4 \ \mu g/cm^2 \ \text{in ethanol} \\ 277.8 \ \mu g/cm^2 \ \text{in cream} & 22.2 \ \mu g/cm^2 \ \text{in ethanol} \\ 1388.9 \ \mu g/cm^2 \ \text{in cream} & 111.1 \ \mu g/cm^2 \ \text{in ethanol} \\ \end{array} $				
$\mu$ g /application/cm <sup>2</sup>	See above				
control substance	Ethanol 96% and glyceryl stearate 15% in water, resp.				
definition of positive	(spotty) erythema of at least 25% of the test area along with homogeneous infiltration or papules regardless of the number				
period	Two weeks for each step until positive reaction or end of study, whichever occurred first				
Results					
PT ED10% (95% CI)	Not calculable; 52 of 60 Patients patch tested positive to 2.5% HICC, 57 / 60 to 5% HICC				
PT ED50% (95% CI)	Not calculable				
PT no effect level (observed)	Not calculable				
ROAT	Cumulative responses:				
	Cream preparation: 2.8 µg/cm <sup>2</sup> : 4.7% 5.6 µg/cm <sup>2</sup> : 12.5% 55.6 µg/cm <sup>2</sup> : 42.2% 277.8 µg/cm <sup>2</sup> : 65.6% 1388.9 µg/cm <sup>2</sup> : 87.5%	Ethanol preparation: 0.2 μg/cm <sup>2</sup> :1.6% 0.4 μg/cm <sup>2</sup> : 3.1% 4.4 μg/cm <sup>2</sup> : 29.7% 22.2 μg/cm <sup>2</sup> : 57.8% 111.1 μg/cm <sup>2</sup> : 82.8%			
Controls	No reactions to vehicle in the patients included into analysis				
Other information	See figure below. Three patients were excluded from the study, so				

results are based on 64 patients.

**Figure 4**: Dose-response curve of 64 patients sensitised to HICC, according to a previous PT, regarding two preparations: perfume and cream, the rhomboid and dot symbol, respectively, indicating the observed response. The curve was fitted by a logistic function (10).



# Isoeugenol

Isoeugenol (11)	
Design	blinded, randomised doses and controlled
Test subjects	20 patients with a positive patch test to isoeugenol
Controls	20 healthy controls
Substance	purity: 98%
Patch test	20 mg solution applied in an 8 mm Finn Chamber occlusion 48 h
-dilution steps	2% to 0.01% (8 steps)
-control/vehicle	petrolatum
-definition of threshold	lowest concentration giving a visible skin reaction in a continuous line
ROAT	outer aspect of upper arms
area	$5 \times 5$ cm (2 areas: one test and one control)
applications/day	two with roll-on
dose	0.2% in ethanol
µg /application/cm <sup>2</sup>	Doses measured to 0.14 -0.13 mg/application the first 14 days = 5.6 $\mu$ g/cm <sup>2</sup>
control substance	ethanol
definition of positive	any degree of reaction
period	Two weeks at upper arm and if negative another two weeks including application to base of neck
Results	
PT ED10% (95% CI)	/
PT ED50% (95% CI)	0.08% 32 μg/cm <sup>2</sup>
PT no effect level (observ)	$< 0.01\% = 0.4 \ \mu g/cm^2$
ROAT	
Dose: 0.2%	12/19 (63%)
Controls	No reactions were seen
Other information	

Isoeugenol (12)	
Design	blinded, randomised
Test subjects	27 patients with a positive patch test to isoeugenol
Controls	20 healthy controls
Substance	purity: 98%
Patch test	15 $\mu$ l solution applied in an 8 mm Finn Chamber occlusion 48 h
-dilution steps	2% to 0.00006% (17 steps)
-control/vehicle	ethanol
-definition of threshold	lowest concentration giving a visible skin reaction in a continuous line of reactions to higher concentrations
ROAT	volar aspect of lower arm
area	3 x 3 cm (2 areas)
applications/day	two with droplet bottle (30 mg per application)
dose	0.05% in ethanol and 0.2%
µg /application/cm <sup>2</sup>	Doses were calculated as mean 2.2 $\mu g/cm^2$ (low conc.) and 9 $\mu g/cm^2$ (high conc.)
control substance	ethanol
definition of positive	clear visible erythema
period	28 days
Results	
PT ED10% (95% CI)	/
PT ED50% (95% CI)	/
PT no effect level (observed)	< 0.0005% (5 ppm)
ROAT	Cumulative responses
Dose 1: 0.05%	10/24 (42%)
Dose 2: 0.2%	16/24 (67%)
Controls	No reactions were seen
Other information	Response to the low concentration in the ROAT appeared after median 15 days and to the high concentration after median 7 days.

Isoeugenol (13)		
Design	blinded, randomised and controlled	
Test subjects	13 patients with a positive patch test to isoeugenol and 4 in part 1 (pre-test)	
Controls	10 healthy controls (dermatitis patients)	
Substance	purity: /	
Patch test	15 $\mu I$ solution applied in an 8 mm Finn Chamber occlusion 48 h	
-dilution steps	2% to 0.00006% (w/v) (16 steps)	
-control/vehicle	ethanol	
-definition of threshold	lowest concentration elicitating at least + reaction	
ROAT	Axilla	
area	10 x 10 cm <sup>2</sup> (estimated)	
applications/day	two with roll-on deodorant (117-586 mg per application of solution) average cases: 266 mg/application controls: only range given	
dose	Part 1: Step 1:0.02% Step 2: 0.063% Step 3:0.2% Part 2: Step1:0.0063% Step 2:0.02% Step 3: 0.063%	
dose/application/cm <sup>2</sup>	Part 2: Step 1: 0.167 Step 2: 0.53 Step 3: 1.67 $\mu$ g/application/ cm <sup>2</sup> (calculated based on data)	
control substance	deodorant matrix	
definition of positive	eczematous response covering 25% of test area	
period	Part one: one week with each concentration: maximum three weeks Part two: two weeks with each concentration: maximum six weeks	
Results		
PT ED10% (95% CI)	/	
PT ED50% (95% CI)	/	
PT no effect level (observed)	<0.0005% (0.15 µg/cm <sup>2</sup> )	
ROAT		
Step 1 (0.0063%)	3/13 (23%)	
Step 2 (0.02%)	9/13 (69%)	
Step 3 (0.063%)	10/13 (77%)	
Controls	No reactions were seen	
Other information	Deodorants containing cinnamal were responsible for all reactions in cinnamal sensitized individuals ( $p$ <0.001) and all control persons were negative ( $p$ <0.001)	

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