



Strada del Petriccio e Belriguardo 35, Siena 53100
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Certificate of Analysis # 2025-1792/EXT/HPLC

Informations provided by the client

Sample Name: CBD OIL 20%
Matrix: Oil
Product: Oil

Laboratory Information

Acceptance Date: 03/09/2025
Test Start Date: 04/09/2025
Sampling: Sample delivered by the customer
Sample ID: 00309251650/EXT

Prepared for

CANNA HEALTH AMSTERDAM
28b Nieuwe Nieuwstraat Amsterdam
VAT NL002505280B09

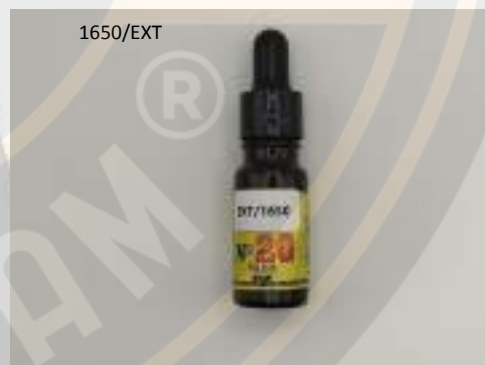
Compound	Result	UM	LOQ	Result (mg/g)	Measurement uncertain
CBDV	0,14	%(m/m)	0,02	1,4	
CBDA	< LOQ	%(m/m)	0,033	< LOQ	
CBGA	< LOQ	%(m/m)	0,035	< LOQ	
CBG	< LOQ	%(m/m)	0,018	< LOQ	
CBD	20,79	%(m/m)	0,015	207,9	
THCV	< LOQ	%(m/m)	0,02	< LOQ	
CBN	< LOQ	%(m/m)	0,02	< LOQ	
d9-THC	< LOQ	%(m/m)	0,028	< LOQ	
CBC	< LOQ	%(m/m)	0,017	< LOQ	
THCA	< LOQ	%(m/m)	0,02	< LOQ	
TOTAL CBD	20,79	%(m/m)	0,02	207,9	
TOTAL CBG	< LOQ	%(m/m)	0,02	< LOQ	
TOTAL THC	< LOQ	%(m/m)	0,02	< LOQ	
TOTAL MOISTURE	NR	%	0,05	NR	

%(m/m) = (Mass of the analyte/Mass of the product as it is)
NR Not Detected
LOD Limit of detection
LOQ Limit of quantification
<LOQ Below the limit of quantification

IST01 REV03 2022 (HPLC)
IST16 REV02 2022 (Termogravimetria)

I cannabinoidi totali vengono calcolati utilizzando le seguenti formule per calcolare la perdita del gruppo carbossilico durante la decarbossilazione
THC TOTALE = THC + (THCA * 0,877)
CBD TOTALE = CBD + (CBDA * 0,877)
CBG TOTALE = CBG + (CBGA * 0,877)

1650/EXT



End of Test Report # Certificate of Analysis # 2025-1792/EXT/HPLC

Test end date: 08/09/2025
Issuing date: 12/09/2025

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Certificate of Analysis # 2025-26/SUBIT/SPEST

Informations provided by the client

Sample Name: CBD OIL 20%
Matrix: Oil
Product: Oil

Laboratory Information

Acceptance Date: 03/09/2025
Test Start Date: 15/09/2025
Sampling: Sample delivered by the customer
Sample ID: 0030925146/SUBIT

Prepared for

CANNA HEALTH AMSTERDAM
28b Nieuwe Nieuwstraat Amsterdam
VAT NL002505280B09

Compound	Result	UM	LOQ	Measurement uncertainty
PCB 77 ^	5,3	pg/g grasso		
PCB 81 ^	< LOQ	pg/g grasso	1	
PCB 105 ^	14,7	pg/g grasso	5	
PCB 114 ^	2,3	pg/g grasso	1	
PCB 118 ^	41,3	pg/g grasso	10	
PCB 123 ^	< LOQ	pg/g grasso	1	
PCB 126 ^	< LOQ	pg/g grasso	0,5	
PCB 156 ^	3,4	pg/g grasso	1	
PCB 157 ^	< LOQ	pg/g grasso	1	
PCB 167 ^	1,7	pg/g grasso	1	
PCB 189 ^	< LOQ	pg/g grasso	1	
PCB 169 ^	< LOQ	pg/g grasso	1	
Sommatoria PCB-DL come WHO-TEQ (upper bound) ^	0,083	pg/g grasso	0,081	
Somma PCDD/F-PCB DL WHO-TEQ (Upper Bound) ^	0,334	pg/g grasso	0,33	
Somma PCDD/F-PCB DL WHO-TEQ (Lower Bound) ^	0,005	pg/g grasso		
Sommatoria PCB-DL as WHO-TEQ (lower bound) ^	0,0024	pg/g grasso		
2,3,7,8-Tetraclorodibenzofurano (TCDF) ^	< LOQ	pg/g grasso	0,02	
1,2,3,4,6,7,8,9-Octaclorodibenzo-p-diossina (OCDD) ^	0,9	pg/g grasso	0,17	
2,3,4,7,8-Pentaclorodibenzofurano (PeCDF) ^	< LOQ	pg/g grasso	0,07	
1,2,3,4,7,8-Esaclorodibenzofurano (ExCDF) ^	< LOQ	pg/g grasso	0,12	
1,2,3,7,8-Pentaclorodibenzofurano (PeCDF) ^	< LOQ	pg/g grasso	0,02	
1,2,3,7,8,9-Esaclorodibenzo-p-diossina (ExCDD) ^	< LOQ	pg/g grasso	0,12	
1,2,3,4,6,7,8-Eptaclorodibenzo-p-diossina (EpCDD) ^	0,28	pg/g grasso	0,12	
1,2,3,6,7,8-Esaclorodibenzofurano (ExCDF) ^	< LOQ	pg/g grasso	0,12	
1,2,3,6,7,8-Esaclorodibenzo-p-diossina (ExCDD) ^	< LOQ	pg/g grasso	0,12	
1,2,3,4,6,7,8,9-Octaclorodibenzofurano (OCDF) ^	< LOQ	pg/g grasso	0,5	
1,2,3,7,8-Pentaclorodibenzo-p-diossina (PeCDD) ^	< LOQ	pg/g grasso	0,07	
1,2,3,4,7,8-Esaclorodibenzo-p-diossina (ExCDD) ^	< LOQ	pg/g grasso	0,12	
1,2,3,4,6,7,8-Eptaclorodibenzofurano (EpCDF) ^	< LOQ	pg/g grasso	0,25	
2,3,4,6,7,8-Esaclorodibenzofurano (ExCDF) ^	< LOQ	pg/g grasso	0,12	
2,3,7,8-Tetraclorodibenzo-p-diossina (TCDD) ^	< LOQ	pg/g grasso	0,06	

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1,2,3,4,7,8,9-Eptaclorodibenzofurano (EpCDF) ^	< LOQ	pg/g grasso	0,25	
1,2,3,7,8,9-Esaclorodibenzofurano (ExCDF) ^	< LOQ	pg/g grasso	0,12	
Sommatoria PCDD, PCDF ^	0,251	pg/g grasso	0,25	
Sommatoria PCDD, PCDF come WHO-TEQ (lower bound) ^	0,003	pg/g grasso		
Benzo[a]antracene ^	< LOQ	µg/kg	0,4	
Benzo[a]pirene ^	< LOQ	µg/kg	0,4	
Indeno[1,2,3-c,d]pirene ^	< LOQ	µg/kg	0,4	
Benzo[b]fluorantene ^	< LOQ	µg/kg	0,4	
Benzo[g,h,i]perilene ^	< LOQ	µg/kg	0,4	
Dibenzo[a,h]antracene ^	< LOQ	µg/kg	0,4	
Benzo[e]pirene ^	0,46	µg/kg	0,4	
Benzo[k]fluorantene ^	< LOQ	µg/kg	0,4	
Crisene ^	< LOQ	µg/kg	0,4	
IPA - somma di: Benzo[a]pirene, Benzo[a]antracene, Benzo[b]fluorantene, Crisene ^	0,46	µg/kg	0,4	
Number of Peroxide ^	6,9	meq O2/kg	0,4	
Parathion-ethyl	< LOQ	mg/kg	0,01	
Chinomethionat	< LOQ	mg/kg	0,01	
trans-Heptachlor epoxide	< LOQ	mg/kg	0,01	
Propoxur	< LOQ	mg/kg	0,01	
Fenhexamid	< LOQ	mg/kg	0,01	
Trichlorfon	< LOQ	mg/kg	0,01	
Oxadiazon	< LOQ	mg/kg	0,01	
Metidathion	< LOQ	mg/kg	0,01	
Carboxin (Carboxin plus its metabolites Carboxin sulfoxide and Oxycarboxin (Carboxin sulfone), expressed as Carboxin)	< LOQ	mg/kg	0,01	
Zoxamide	< LOQ	mg/kg	0,01	
* Oxamyl	< LOQ	mg/kg	0,01	
* Ethofumesate	< LOQ	mg/kg	0,01	
Nitrofen	< LOQ	mg/kg	0,01	
Difenoconazole	< LOQ	mg/kg	0,01	
Metobromuron	< LOQ	mg/kg	0,01	
Cyanazin	< LOQ	mg/kg	0,01	
Tetraconazole	< LOQ	mg/kg	0,01	
Abamectin (Sum of Avermectin B1a, Avermectin B1b and delta-8,9 isomer of Avermectin B1a, expressed as Avermectin B1a)	< LOQ	mg/kg	0,01	
Avermectin B1b	< LOQ	mg/kg	0,01	
Atrazine-desisopropyl	< LOQ	mg/kg	0,01	
Bifenthrin (sum of isomers)	< LOQ	mg/kg	0,01	
Fenoxycarb	< LOQ	mg/kg	0,01	
Phosmet (Sum of Phosmet and Phosmet oxon expressed as Phosmet)	< LOQ	mg/kg	0,01	
Quintozene	< LOQ	mg/kg	0,01	
Chlorpyrifos-methyl	< LOQ	mg/kg	0,01	

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* Famoxadone	< LOQ	mg/kg	0,01	
Dicofol (Sum of p,p' and o,p' isomers)	< LOQ	mg/kg	0,01	
Fonofos	< LOQ	mg/kg	0,01	
Oxyfluorfen	< LOQ	mg/kg	0,01	
cis-Heptachloroepoxide	< LOQ	mg/kg	0,01	
TFNA	< LOQ	mg/kg	0,01	
Fenthion-oxon-sulfone	< LOQ	mg/kg	0,01	
Piperonyl butoxide	< LOQ	mg/kg	0,01	
Flucythrinate (Flucythrinate including other mixtures of constituent isomers (Sum of isomers))	< LOQ	mg/kg	0,01	
Ametryn	< LOQ	mg/kg	0,01	
Clomazone	< LOQ	mg/kg	0,01	
Fenvalerate (any ratio of constituent isomers (RR, SS, RS & SR) including Esfenvalerate)	< LOQ	mg/kg	0,01	
TFNG	< LOQ	mg/kg	0,01	
2,4-Dimethylaniline [2,4 DMA]	< LOQ	mg/kg	0,01	
Pyraclostrobin	< LOQ	mg/kg	0,01	
Triflumizole (Triflumizole and metabolite FM-6-1(N-(4-chloro-2-Trifluoromethylphenyl)-n-Propoxyacetamide), expressed as Triflumizole)	< LOQ	mg/kg	0,01	
Diclofop	< LOQ	mg/kg	0,01	
* Oxydemeton-methyl (Sum of Oxydemeton-methyl and Demeton-S-methylsulfone expressed as Oxydemeton-methyl)	< LOQ	mg/kg	0,01	
Chlorotoluron	< LOQ	mg/kg	0,01	
* Ethofumesate (Sum of ethofumesate, 2-keto-ethofumesate, open-ring-2-keto-ethofumesate and its conjugate, expressed as ethofumesate)	< LOQ	mg/kg	0,01	
Nitrothal-isopropyl	< LOQ	mg/kg	0,01	
* Spinosyn A	< LOQ	mg/kg	0,01	
Pymetrozine	< LOQ	mg/kg	0,01	
Benfuracarb	< LOQ	mg/kg	0,01	
Phosmet	< LOQ	mg/kg	0,01	
Fenamiphos-sulfoxide	< LOQ	mg/kg	0,01	
N-2,4-Dimethylphenyl-N'-methylformamidine [DMPF]	< LOQ	mg/kg	0,01	
Tridemorph	< LOQ	mg/kg	0,01	
Fipronil (Sum of Fipronil and Fipronil Sulfone expressed as Fipronil)	< LOQ	mg/kg	0,01	
Lenacil	< LOQ	mg/kg	0,01	
Pyridaphenthion	< LOQ	mg/kg	0,01	
Fenthion (Sum)	< LOQ	mg/kg	0,01	
* Fenuron	< LOQ	mg/kg	0,01	
Captan (Sum of Captan and Tetrahydrophthalimide exp as Captan)	< LOQ	mg/kg	0,01	
Dimethomorph (Sum of isomers)	< LOQ	mg/kg	0,01	
Fenpyroximate	< LOQ	mg/kg	0,01	

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Cadusafos	< LOQ	mg/kg	0,01	
Bendiocarb	< LOQ	mg/kg	0,01	
Chlorpyrifos-ethyl	< LOQ	mg/kg	0,01	
Fenitrothion	< LOQ	mg/kg	0,01	
Metazachlor	< LOQ	mg/kg	0,01	
* Tebufenozide	< LOQ	mg/kg	0,01	
Cymiazole	< LOQ	mg/kg	0,01	
Bromophos-ethyl	< LOQ	mg/kg	0,01	
Mandipropamid (any ratio of constituent isomers)	< LOQ	mg/kg	0,01	
Carboxin	< LOQ	mg/kg	0,01	
Avermectin B1a	< LOQ	mg/kg	0,01	
Amitraz	< LOQ	mg/kg	0,01	
Cyfluthrin-beta	< LOQ	mg/kg	0,01	
Imazalil (any ratio of constituent isomers)	< LOQ	mg/kg	0,01	
Phosmet oxon	< LOQ	mg/kg	0,01	
Iodofenphos	< LOQ	mg/kg	0,01	
Tebupirimfos	< LOQ	mg/kg	0,01	
alpha-HCH	< LOQ	mg/kg	0,01	
Tribenuron-methyl	< LOQ	mg/kg	0,01	
Fuberidazole	< LOQ	mg/kg	0,01	
Bromocyclen	< LOQ	mg/kg	0,01	
Iprodione	< LOQ	mg/kg	0,01	
Fludioxonil	< LOQ	mg/kg	0,01	
Paclobutrazol (Sum of constituent isomers)	< LOQ	mg/kg	0,01	
Atrazine-desethyl	< LOQ	mg/kg	0,01	
Atrazine	< LOQ	mg/kg	0,01	
Chlorfenson	< LOQ	mg/kg	0,01	
Flufenacet-ethane sulfonic acid (ESA)	< LOQ	mg/kg	0,01	
Metribuzin	< LOQ	mg/kg	0,01	
Fenpropimorph (sum of isomers)	< LOQ	mg/kg	0,01	
Parathion-methyl	< LOQ	mg/kg	0,01	
Flonicamid (Sum of Flonicamid and TFNA, TFNG expressed as Flonicamid)	< LOQ	mg/kg	0,01	
Pyrazophos	< LOQ	mg/kg	0,01	
Fenarimol	< LOQ	mg/kg	0,01	
Tebuconazole	< LOQ	mg/kg	0,01	
Vamidothion	< LOQ	mg/kg	0,01	
Prochloraz	< LOQ	mg/kg	0,01	
Formothion	< LOQ	mg/kg	0,01	
Promecarb	< LOQ	mg/kg	0,01	
Terbutylazine-desethyl	< LOQ	mg/kg	0,01	
Tetrahydrophthalimide	< LOQ	mg/kg	0,01	
Carfentrazone-ethyl (Carfentrazone free acid expressed as Carfentrazone-ethyl)	< LOQ	mg/kg	0,01	
Fenthion-sulfone	< LOQ	mg/kg	0,01	

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Methiocarb	< LOQ	mg/kg	0,01	
Phorate-oxon-sulfone	< LOQ	mg/kg	0,01	
Isofenphos	< LOQ	mg/kg	0,01	
FM-6	< LOQ	mg/kg	0,01	
Hexythiazox	< LOQ	mg/kg	0,01	
Cyprodinil	0,031	mg/kg	0,01	
Disulfoton	< LOQ	mg/kg	0,01	
Alachlor	< LOQ	mg/kg	0,01	
Methoxychlor	< LOQ	mg/kg	0,01	
Terbuthylazine	< LOQ	mg/kg	0,01	
Azinphos-ethyl	< LOQ	mg/kg	0,01	
Captan	< LOQ	mg/kg	0,01	
Pyridaben	< LOQ	mg/kg	0,01	
Dichlofluanid	< LOQ	mg/kg	0,01	
Isoxaflutole-diketonitrile	< LOQ	mg/kg	0,01	
Flufenacet	< LOQ	mg/kg	0,01	
Propham	< LOQ	mg/kg	0,01	
Pirimiphos-ethyl	< LOQ	mg/kg	0,01	
Dieldrin (Sum of Dieldrin and Aldrin expressed as Dieldrin)	< LOQ	mg/kg	0,01	
Disulfoton (Sum of Disulfoton, Disulfoton-sulfone, Disulfoton-sulfoxide expressed as Disulfoton)	< LOQ	mg/kg	0,01	
Acrinathrin	< LOQ	mg/kg	0,01	
Quizalofop acid	< LOQ	mg/kg	0,01	
Endrin	< LOQ	mg/kg	0,01	
Methacrifos	< LOQ	mg/kg	0,01	
Spinosad (Spinosad, Sum of Spinosyn A and Spinosyn D)	< LOQ	mg/kg	0,01	
Carbosulfan	< LOQ	mg/kg	0,01	
Furathiocarb	< LOQ	mg/kg	0,01	
Azinphos-methyl	< LOQ	mg/kg	0,01	
Nuarimol	< LOQ	mg/kg	0,01	
Dicloran	< LOQ	mg/kg	0,01	
Flutriafof	< LOQ	mg/kg	0,01	
Flubenzimine	< LOQ	mg/kg	0,01	
Malaoxon	< LOQ	mg/kg	0,01	
Linuron	< LOQ	mg/kg	0,01	
Flurochloridone (Sum of cis- and trans- isomers)	< LOQ	mg/kg	0,01	
Prometon	< LOQ	mg/kg	0,01	
Benzoximate MP/C/22 rev 7 2020	< LOQ	mg/kg	0,01	
Buprofezin	< LOQ	mg/kg	0,01	
Carbophenothion	< LOQ	mg/kg	0,01	
DDT (Sum of p,p'-DDT, o,p'-DDT, p,p'-DDE and p,p'-TDE (DDD) expressed as DDT)	< LOQ	mg/kg	0,01	
p,p'-DDT	< LOQ	mg/kg	0,01	
Methamidophos	< LOQ	mg/kg	0,01	
* Metoxuron	< LOQ	mg/kg	0,01	

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Phorate	< LOQ	mg/kg	0,01	
Chlorobenzilate	< LOQ	mg/kg	0,01	
Amitraz (included metabolite containing 2,4-DMA expressed as Amitraz)	< LOQ	mg/kg	0,01	
Flusilazole	< LOQ	mg/kg	0,01	
Trifluralin	< LOQ	mg/kg	0,01	
Propaquizafop	< LOQ	mg/kg	0,01	
BTS 44595	< LOQ	mg/kg	0,01	
Heptachlor	< LOQ	mg/kg	0,01	
Fenothiocarb	< LOQ	mg/kg	0,01	
Diniconazole (Sum of isomers)	< LOQ	mg/kg	0,01	
Flufenacet (Sum of all compounds containing the N fluorophenyl-N-isopropyl moiety expressed as Flufenacet equivalent)	< LOQ	mg/kg	0,01	
Dichlobenil	< LOQ	mg/kg	0,01	
Pirimiphos-methyl	< LOQ	mg/kg	0,01	
Propachlor	< LOQ	mg/kg	0,01	
Naled	< LOQ	mg/kg	0,01	
Prothioconazole	< LOQ	mg/kg	0,01	
beta-HCH	< LOQ	mg/kg	0,01	
Fipronil-sulfone	< LOQ	mg/kg	0,01	
Chlorothalonil	< LOQ	mg/kg	0,01	
Alphamethrin	< LOQ	mg/kg	0,01	
Metazachlor ESA (479M08)	< LOQ	mg/kg	0,01	
Fenpropidin (Sum of Fenpropidin and its salts, expressed as Fenpropidin)	< LOQ	mg/kg	0,01	
Aldrin	< LOQ	mg/kg	0,01	
Tebufenpyrad	< LOQ	mg/kg	0,01	
Bitertanol (sum of isomers)	< LOQ	mg/kg	0,01	
Terbumeton	< LOQ	mg/kg	0,01	
Diclobutrazol	< LOQ	mg/kg	0,01	
Carboxin sulfoxide	< LOQ	mg/kg	0,01	
Cyproconazole	< LOQ	mg/kg	0,01	
Paraoxon-methyl	< LOQ	mg/kg	0,01	
Chlordane (Sum of cis-Chlordane and trans-Chlordane)	< LOQ	mg/kg	0,01	
* Ethiofencarb	< LOQ	mg/kg	0,01	
Allethrin	< LOQ	mg/kg	0,01	
o-p'-DDT	< LOQ	mg/kg	0,01	
Butoxycarboxim	< LOQ	mg/kg	0,01	
Cyazofamid	< LOQ	mg/kg	0,01	
Triflusalufuron (6-(2,2,2-trifluoroethoxy)-1,3,5-triazine-2,4-diamine (IN-M7222))	< LOQ	mg/kg	0,01	
Flurprimidol	< LOQ	mg/kg	0,01	
Carfentrazone acid	< LOQ	mg/kg	0,01	

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Phosalone	< LOQ	mg/kg	0,01	
Biphenyl	< LOQ	mg/kg	0,01	
Isoproturon	< LOQ	mg/kg	0,01	
Fenthion-oxon	< LOQ	mg/kg	0,01	
Cypermethrin (Sum of isomers)	< LOQ	mg/kg	0,01	
Fluiconazole	< LOQ	mg/kg	0,01	
Benomyl	< LOQ	mg/kg	0,01	
Furalaxyl	< LOQ	mg/kg	0,01	
N-2,4-Dimethylphenyl-formamide [DMF]	< LOQ	mg/kg	0,01	
* Rimsulfuron	< LOQ	mg/kg	0,01	
Carfentrazone-ethyl	< LOQ	mg/kg	0,01	
Aldicarb-sulfoxide	< LOQ	mg/kg	0,01	
Fenthion-oxon-sulfoxide	< LOQ	mg/kg	0,01	
Carbaryl	< LOQ	mg/kg	0,01	
Chlorthal-dimethyl	< LOQ	mg/kg	0,01	
Triazophos	< LOQ	mg/kg	0,01	
Mecarbam	< LOQ	mg/kg	0,01	
Aclonifen	< LOQ	mg/kg	0,01	
Chlorfenapyr	< LOQ	mg/kg	0,01	
Mevinphos (Sum of isomer E and Z)	< LOQ	mg/kg	0,01	
Phthalimide	< LOQ	mg/kg	0,01	
trans-Chlordane	< LOQ	mg/kg	0,01	
Tetradifon	< LOQ	mg/kg	0,01	
2-keto-ethofumesate	< LOQ	mg/kg	0,01	
Diclofop-methyl	< LOQ	mg/kg	0,01	
Thiacloprid	< LOQ	mg/kg	0,01	
Oxycarboxin	< LOQ	mg/kg	0,01	
Dinitramine	< LOQ	mg/kg	0,01	
Disulfoton-sulfoxyde	< LOQ	mg/kg	0,01	
Chloropropylate	< LOQ	mg/kg	0,01	
Simazine	< LOQ	mg/kg	0,01	
Clethodim	< LOQ	mg/kg	0,01	
Phenmedipham	< LOQ	mg/kg	0,01	
Penconazole (Sum of constituent isomers)	< LOQ	mg/kg	0,01	
* Ethiofencarb-sulfoxide	< LOQ	mg/kg	0,01	
cis-Chlordane	< LOQ	mg/kg	0,01	
4-Phenylphenol	< LOQ	mg/kg	0,01	
Isofenphos-methyl	< LOQ	mg/kg	0,01	
Flufenacet alcohol	< LOQ	mg/kg	0,01	
Terbutryn	< LOQ	mg/kg	0,01	
* Etofenprox	< LOQ	mg/kg	0,01	
Isoxaflutole (Sum of Isoxaflutole and its diketonitrile-metabolite, expressed as Isoxaflutole)	< LOQ	mg/kg	0,01	
Fenson	< LOQ	mg/kg	0,01	

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Certificate of Analysis # 2025-26/SUBIT/SPEST

Pencycuron	< LOQ	mg/kg	0,01	
Benfluralin	< LOQ	mg/kg	0,01	
delta-HCH	< LOQ	mg/kg	0,01	
Parathion-methyl (Sum of Parathion-methyl and Paraoxon-methyl expressed as Parathion-methyl)	< LOQ	mg/kg	0,01	
Coumatetralyl	< LOQ	mg/kg	0,01	
Bromopropylate	< LOQ	mg/kg	0,01	
Clothianidin	< LOQ	mg/kg	0,01	
Metazaclo metabolite (479M16)	< LOQ	mg/kg	0,01	
Spiromesifen	< LOQ	mg/kg	0,01	
BifenoX	< LOQ	mg/kg	0,01	
Etaconazole	< LOQ	mg/kg	0,01	
Prothoate	< LOQ	mg/kg	0,01	
Disulfoton-sulfone	< LOQ	mg/kg	0,01	
Ethoprophos	< LOQ	mg/kg	0,01	
Isopropalin	< LOQ	mg/kg	0,01	
Dichlorvos	< LOQ	mg/kg	0,01	
Etrimfos	< LOQ	mg/kg	0,01	
Fenamiphos	< LOQ	mg/kg	0,01	
Trifloxystrobin	< LOQ	mg/kg	0,01	
Dialifos	< LOQ	mg/kg	0,01	
Fluazinam	< LOQ	mg/kg	0,01	
Diethofencarb	< LOQ	mg/kg	0,01	
Phorate-oxon	< LOQ	mg/kg	0,01	
Quizalofop (Sum of Quizalofop, its salts, its esters (including Propaquizafop) and its conjugates, expressed as Quizalofop (any ratio of constituent isomers))	< LOQ	mg/kg	0,01	
Monolinuron	< LOQ	mg/kg	0,01	
Diclofop (Sum Diclofop-methyl and Diclofop acid expressed as Diclofop-methyl)	< LOQ	mg/kg	0,01	
Iprobenfos	< LOQ	mg/kg	0,01	
PyrifenoX	< LOQ	mg/kg	0,01	
Iprovalicarb	< LOQ	mg/kg	0,01	
Propiconazole (Sum of isomers)	< LOQ	mg/kg	0,01	
Methiocarb (Sum of Methiocarb, Methiocarb-sulfone, Methiocarb-sulfoxide expressed as Methiocarb)	< LOQ	mg/kg	0,01	
Tefluthrin	< LOQ	mg/kg	0,01	
Fluopicolide	< LOQ	mg/kg	0,01	
Tolyfluanid	< LOQ	mg/kg	0,01	
Metolachlor and S-Metolachlor (Metolachlor including other mixtures of constituent isomers including S-Metolachlor (Sum of isomers))	< LOQ	mg/kg	0,01	
Thiamethoxam	< LOQ	mg/kg	0,01	
Benzoylprop-ethyl	< LOQ	mg/kg	0,01	
Chlorantraniliprole (DPX E-2Y45)	< LOQ	mg/kg	0,01	
Terbufos	< LOQ	mg/kg	0,01	

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Certificate of Analysis # 2025-26/SUBIT/SPEST

Pentachloroaniline	< LOQ	mg/kg	0,01	
2-Phenylphenol (Sum of 2-Phenylphenol and conjugates, expressed as 2-Phenylphenol)	< LOQ	mg/kg	0,01	
p-p'-DDD	< LOQ	mg/kg	0,01	
Fenamiphos-sulfone	< LOQ	mg/kg	0,01	
Spiroxamine (Sum of isomers)	< LOQ	mg/kg	0,01	
Flufenacet thioglycolate sulfoxide	< LOQ	mg/kg	0,01	
Diflufenican	< LOQ	mg/kg	0,01	
Aldicarb (Sum of Aldicarb and Aldicarb-sulfone, Aldicarb-sulfoxide expressed as Aldicarb)	< LOQ	mg/kg	0,01	
Chloroxuron	< LOQ	mg/kg	0,01	
Edifenphos	< LOQ	mg/kg	0,01	
Butocarboxim	< LOQ	mg/kg	0,01	
Aldicarb	< LOQ	mg/kg	0,01	
Acibenzolar-S-methyl (Sum of Acibenzolar-S-methyl and Acibenzolar acid expressed as Acibenzolar-S-methyl)	< LOQ	mg/kg	0,01	
Azadirachtin	< LOQ	mg/kg	0,01	
Captafol	< LOQ	mg/kg	0,01	
Metazaclo OA (479M04)	< LOQ	mg/kg	0,01	
Dichlofenthion	< LOQ	mg/kg	0,01	
Tolyfluanid (Sum of Tolyfluanid and DMST expressed as Tolyfluanid)	< LOQ	mg/kg	0,01	
Flamprop-isopropyl	< LOQ	mg/kg	0,01	
Acibenzolar-S-methyl	< LOQ	mg/kg	0,01	
Quinalphos	< LOQ	mg/kg	0,01	
Imazamox (Sum of Imazamox and its salts, expressed as Imazamox)	< LOQ	mg/kg	0,01	
* Ethiofencarb-sulfone	< LOQ	mg/kg	0,01	
Folpet (Sum of Folpet and Phtalimide expressed as Folpet)	< LOQ	mg/kg	0,01	
Napropamide	< LOQ	mg/kg	0,01	
Malathion (Sum of Malathion and Malaaxon expressed as Malathion)	< LOQ	mg/kg	0,01	
Spinosyn D	< LOQ	mg/kg	0,01	
Benalaxyl including other mixtures of constituent isomers including benalaxyl-M (sum of isomers)	< LOQ	mg/kg	0,01	
Pendimethalin	< LOQ	mg/kg	0,01	
Propachlor oxalinic acid	< LOQ	mg/kg	0,01	
Carbofuran-3-hydroxy	< LOQ	mg/kg	0,01	
Avermectin B1a 8,9z	< LOQ	mg/kg	0,01	
Propachlor: oxalinic derivate of Propachlor, expressed as Propachlor	< LOQ	mg/kg	0,01	
Methiocarb-sulfone	< LOQ	mg/kg	0,01	
Prothioconazole-desthio	0,054	mg/kg	0,01	
Chlorfenvinphos	< LOQ	mg/kg	0,01	
Cycloxydim	< LOQ	mg/kg	0,01	

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Folpet	< LOQ	mg/kg	0,01	
Fluazifop-P (Sum of all the constituent isomers of Fluazifop, its esters and its conjugates, expressed as Fluazifop)	< LOQ	mg/kg	0,01	
Cymoxanil	< LOQ	mg/kg	0,01	
alpha-Endosulfan	< LOQ	mg/kg	0,01	
Endrin aldehyde	< LOQ	mg/kg	0,01	
Methiocarb-sulfoxide	< LOQ	mg/kg	0,01	
Mepanipyrim	< LOQ	mg/kg	0,01	
Pirimicarb	< LOQ	mg/kg	0,01	
Myclobutanil	< LOQ	mg/kg	0,01	
Chlormephos	< LOQ	mg/kg	0,01	
Metazachlor (Sum of metabolites 479M04, 479M08 and 479M16, expressed as Metazachlor)	< LOQ	mg/kg	0,01	
Thiophanate-methyl	< LOQ	mg/kg	0,01	
Permethrin (Sum of isomers)	< LOQ	mg/kg	0,01	
Fipronil	< LOQ	mg/kg	0,01	
Monocrotophos	< LOQ	mg/kg	0,01	
Chloridazon-desphenyl	< LOQ	mg/kg	0,01	
Propyzamide	< LOQ	mg/kg	0,01	
Spirodiclofen	< LOQ	mg/kg	0,01	
Bupirimate	< LOQ	mg/kg	0,01	
Kresoxim-methyl	< LOQ	mg/kg	0,01	
Chlorpropham	< LOQ	mg/kg	0,01	
* Methoxyfenozide	< LOQ	mg/kg	0,01	
Diuron	< LOQ	mg/kg	0,01	
Chlozolinate	< LOQ	mg/kg	0,01	
Propamocarb (Sum of Propamocarb and its salts, expressed as Propamocarb)	0,027	mg/kg	0,01	
Tetramethrin	< LOQ	mg/kg	0,01	
Hexaconazole	< LOQ	mg/kg	0,01	
epsilon-HCH	< LOQ	mg/kg	0,01	
Fenpropathrin	< LOQ	mg/kg	0,01	
* Clethodim (Sum of Sethoxydim and Clethodim including degradation products calculated as Sethoxydim)	< LOQ	mg/kg	0,01	
Prothiophos	< LOQ	mg/kg	0,01	
gamma HCH [Lindane]	< LOQ	mg/kg	0,01	
Ethalfuralin	< LOQ	mg/kg	0,01	
Endosulfan (Sum of Alpha and Beta and Sulfate expressed as Endosulfan)	< LOQ	mg/kg	0,01	
Fenthion-sulfoxide	< LOQ	mg/kg	0,01	
Flonicamid	< LOQ	mg/kg	0,01	
Fenazaquin	< LOQ	mg/kg	0,01	
Pyrethrins	< LOQ	mg/kg	0,01	
Metalaxyl and Metalaxyl-M (Metalaxyl including other mixtures of constituent isomers including Metalaxyl-M (Sum of isomers))	< LOQ	mg/kg	0,01	

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Certificate of Analysis # 2025-26/SUBIT/SPEST

HCH (Hexachlorocyclohexane) (Sum of isomers Alpha, Beta, Delta and Epsilon)	< LOQ	mg/kg	0,01	
Tolclofos-methyl	< LOQ	mg/kg	0,01	
Azaconazole	< LOQ	mg/kg	0,01	
Chlorthiamid	< LOQ	mg/kg	0,01	
Isoxaben	< LOQ	mg/kg	0,01	
Epoxiconazole	< LOQ	mg/kg	0,01	
Hexachlorobenzene	< LOQ	mg/kg	0,01	
Perthane	< LOQ	mg/kg	0,01	
Dimethoate	< LOQ	mg/kg	0,01	
Fenamiphos (Sum of Fenamiphos and Fenamiphos-sulfone, Fenamiphos-sulfoxide expressed as Fenamiphos)	< LOQ	mg/kg	0,01	
Quintozene (Sum of Quintozene and Pentachloroaniline expressed as Quintozene)	< LOQ	mg/kg	0,01	
Fenthion	< LOQ	mg/kg	0,01	
Procymidone	< LOQ	mg/kg	0,01	
Chloridazon (sum of Chloridazon and Chloridazon-desphenyl, expressed as Chloridazon)	< LOQ	mg/kg	0,01	
Cyfluthrin (Sum of isomers)	< LOQ	mg/kg	0,01	
Heptachlor (Sum of Heptachlor and Heptachlor epoxide expressed as Heptachlor)	< LOQ	mg/kg	0,01	
Bromophos-methyl	< LOQ	mg/kg	0,01	
Anilazine	< LOQ	mg/kg	0,01	
Vinclozolin	< LOQ	mg/kg	0,01	
Dieldrin	< LOQ	mg/kg	0,01	
Tralkoxydim (sum of the constituent isomers of tralkoxydim)	< LOQ	mg/kg	0,01	
BTS 44596	< LOQ	mg/kg	0,01	
* Metaflumizone (Sum of isomer E and Z)	< LOQ	mg/kg	0,01	
Paraoxon-ethyl	< LOQ	mg/kg	0,01	
Carbendazim	< LOQ	mg/kg	0,01	
* Demeton-S-methyl-sulfone	< LOQ	mg/kg	0,01	
* Milbemectin (Sum of Milbemycin A4 and A3 expressed as Milbemectin)	< LOQ	mg/kg	0,01	
Phorate (sum of Phorate, its oxygen analogue and their sulfones expressed as Phorate)	< LOQ	mg/kg	0,01	
Tetrachlorvinphos	< LOQ	mg/kg	0,01	
Deltamethrin	< LOQ	mg/kg	0,01	
Malathion	< LOQ	mg/kg	0,01	
Phosphamidon	< LOQ	mg/kg	0,01	
Endosulfan-sulfate	< LOQ	mg/kg	0,01	
Dicofolophos	< LOQ	mg/kg	0,01	
Diazinon	< LOQ	mg/kg	0,01	
Pyrimethanil	< LOQ	mg/kg	0,01	
* Fenamidone	< LOQ	mg/kg	0,01	
Propazine	< LOQ	mg/kg	0,01	
Imidacloprid	< LOQ	mg/kg	0,01	

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Ethion	< LOQ	mg/kg	0,01	
Omethoate	< LOQ	mg/kg	0,01	
Flucycloxuron	< LOQ	mg/kg	0,01	
Carbofuran	< LOQ	mg/kg	0,01	
Triflumizole	< LOQ	mg/kg	0,01	
Phorate-sulfone	< LOQ	mg/kg	0,01	
4-Fluoro-N-isopropylaniline	< LOQ	mg/kg	0,01	
* Fluoxastrobin (sum of fluoxastrobin and its Z-isomer)	< LOQ	mg/kg	0,01	
Cypermethrin	< LOQ	mg/kg	0,01	
Benomyl (Sum of Benomyl and Carbendazim espressed as Carbendazim)	< LOQ	mg/kg	0,01	
Coumaphos	< LOQ	mg/kg	0,01	
Pyriproxyfen	< LOQ	mg/kg	0,01	
Methomyl	< LOQ	mg/kg	0,01	
* Nicosulfuron	< LOQ	mg/kg	0,01	
Acibenzolar acid	< LOQ	mg/kg	0,01	
Metamitron	< LOQ	mg/kg	0,01	
Fenbuconazole (Sum of constituent enantiomers)	< LOQ	mg/kg	0,01	
Chloridazon	< LOQ	mg/kg	0,01	
Phenthoate	< LOQ	mg/kg	0,01	
p-p'-DDE	< LOQ	mg/kg	0,01	
Heptenophos	< LOQ	mg/kg	0,01	
Propanil	< LOQ	mg/kg	0,01	
Tricyclazole	< LOQ	mg/kg	0,01	
Sulfotep	< LOQ	mg/kg	0,01	
lambda-Cyhalothrin (includes gamma-Cyhalothrin) (sum of R,S and S,R isomers)	< LOQ	mg/kg	0,01	
Prochloraz (Sum of Prochloraz, BTS 44595 (M201-04), BTS 44596 (M201-03), expressed as Prochloraz)	< LOQ	mg/kg	0,01	
tau-Fluvalinate	< LOQ	mg/kg	0,01	
Fosthiazate	< LOQ	mg/kg	0,01	
* Rotenone	< LOQ	mg/kg	0,01	
Profenofos	< LOQ	mg/kg	0,01	
Quinoxifen	< LOQ	mg/kg	0,01	
Bromuconazole (sum of diastereoisomers)	< LOQ	mg/kg	0,01	
Pethoxamid	< LOQ	mg/kg	0,01	
Acetamiprid	< LOQ	mg/kg	0,01	
Bifenazate	< LOQ	mg/kg	0,01	
Boscalid	< LOQ	mg/kg	0,01	
Fenpyrazamine	< LOQ	mg/kg	0,01	
Fluopyram	< LOQ	mg/kg	0,01	
DEET Diethyl-m-toluamid,N,N	< LOQ	mg/kg	0,01	
Azoxystrobin	< LOQ	mg/kg	0,01	
Fluxapyroxad	< LOQ	mg/kg	0,01	

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Emamectin B1a and its salts, expressed as emamectin B1a (free base)	< LOQ	mg/kg	0,01	
Prosulfocarb	< LOQ	mg/kg	0,01	

%(m/m) = (Mass of the analyte/Mass of the product as it is)
NR Not Detected
LOD Limit of detection
LOQ Limit of quantification
<LOQ Below the limit of quantification

^ Subappalto
Metodi:
UNI EN 15662:2018

146/SUBIT



End of Test Report # Certificate of Analysis # 2025-26/SUBIT/SPEST

Test end date: 25/09/2025
Issuing date: 06/10/2025

The results reported in the Test Report refer only to the sample actually tested, as received. The Laboratory declines all responsibility for the methods of carrying out sampling, transport and storage of the samples until delivery, if carried out by the customer. The Laboratory assumes responsibility for the information and data contained in the test report, with the exclusion of what is declared by the customer. This CoA can only be reproduced in complete form: if not complete, it is allowed only after written authorization from the Laboratory Manager. Copies of this Test Report and related documents are kept for 4 years. The laboratory declines responsibility for any information provided by the customer that may affect the validity of the results. Test Report digitally signed and compliant pursuant to art. 23 Legislative Decree 7 March 2005 No. 82 CAD and subsequent amendments and additions.



Certificate of Analysis # 2025-14/SUBIT/SRSOLV

Informations provided by the client

Sample Name: CBD OIL 20%
Matrix: Oil
Product: Oil

Laboratory Information

Acceptance Date: 03/09/2025
Test Start Date: 26/09/2025
Sampling: Sample delivered by the customer
Sample ID: 0030925147/SUBIT

Prepared for

CANNA HEALTH AMSTERDAM
28b Nieuwe Nieuwstraat Amsterdam
VAT NL002505280B09

Compound	Result	UM	LOQ	Measurement uncertainty
Toluene ^	<3	mg/kg	3	
Butyl acetate ^	< LOQ	mg/kg	2	
Methyl-1-propanol ^	< LOQ	mg/kg	1	
Dichloromethane ^	< LOQ	mg/kg	3	
Cyclohexane ^	4,6	mg/kg	1	
Hexane ^	60	mg/kg	1	
Propan-2-olo ^	<3	mg/kg	3	
MEK (Methyl ethyl chetone) ^	< LOQ	mg/kg	1	
Butan-1-olo ^	< LOQ	mg/kg	1	
Propanol ^	<1	mg/kg	1	
Chloroform (trichloromethane) ^	< LOQ	mg/kg	3	
Methanol ^	< LOQ	mg/kg	3	
Diethyl ether ^	< LOQ	mg/kg	1	
Methyl acetate ^	< LOQ	mg/kg	1	
Acetone ^	< LOQ	mg/kg	3	
Benzene ^	< LOQ	mg/kg	3	
Butan-2-olo ^	< LOQ	mg/kg	1	
Ethyl acetate ^	< LOQ	mg/kg	1	
Ethanol ^	3,15	mg/kg	1	

%(m/m) = (Mass of the analyte/Mass of the product as it is)
NR Not Detected
LOD Limit of detection
LOQ Limit of quantification
<LOQ Below the limit of quantification

^ Subappalto
Metodi: MP/C/827 rev 0 2013

147/SUBIT



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Certificate of Analysis # 2025-14/SUBIT/SRSOLV

End of Test Report # Certificate of Analysis # 2025-14/SUBIT/SRSOLV

Test end date: 27/09/2025

Issuing date: 06/10/2025

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The results reported in the Test Report refer only to the sample actually tested, as received. The Laboratory declines all responsibility for the methods of carrying out sampling, transport and storage of the samples until delivery, if carried out by the customer. The Laboratory assumes responsibility for the information and data contained in the test report, with the exclusion of what is declared by the customer. This CoA can only be reproduced in complete form: if not complete, it is allowed only after written authorization from the Laboratory Manager. Copies of this Test Report and related documents are kept for 4 years. The laboratory declines responsibility for any information provided by the customer that may affect the validity of the results.

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Certificate of Analysis # 2025-12/SUBIT/SMTOX

Informations provided by the client

Sample Name: CBD OIL 20%
Matrix: Oil
Product: Oil

Laboratory Information

Acceptance Date: 03/09/2025
Test Start Date: 16/09/2025
Sampling: Sample delivered by the customer
Sample ID: 0030925148/SUBIT

Prepared for

CANNA HEALTH AMSTERDAM
28b Nieuwe Nieuwstraat Amsterdam
VAT NL002505280B09

Compound	Result	UM	LOQ	Measurement uncertainty
Aflatoxin B2 MS ^	< LOQ	µg/kg	0,5	
Aflatoxin B1 MS ^	< LOQ	µg/kg	0,5	
Aflatoxin G1 MS ^	< LOQ	µg/kg	0,5	
Aflatoxin G2 MS ^	< LOQ	µg/kg	0,5	
Ochratoxin (OTA) MS ^	< LOQ	µg/kg	0,3	

%(m/m) = (Mass of the analyte/Mass of the product as it is)
NR Not Detected
LOD Limit of detection
LOQ Limit of quantification
<LOQ Below the limit of quantification

^ Subappalto
Metodi: MP/C/16 rev 11 2019

148/SUBIT



End of Test Report # Certificate of Analysis # 2025-12/SUBIT/SMTOX

Test end date: 17/09/2025
Issuing date: 30/09/2025

The results reported in the Test Report refer only to the sample actually tested, as received. The Laboratory declines all responsibility for the methods of carrying out sampling, transport and storage of the samples until delivery, if carried out by the customer. The Laboratory assumes responsibility for the information and data contained in the test report, with the exclusion of what is declared by the customer. This CoA can only be reproduced in complete form: if not complete, it is allowed only after written authorization from the Laboratory Manager. Copies of this Test Report and related documents are kept for 4 years. The laboratory declines responsibility for any information provided by the customer that may affect the validity of the results. Test Report digitally signed and compliant pursuant to art. 23 Legislative Decree 7 March 2005 No. 82 CAD and subsequent amendments and additions.



Certificate of Analysis # 2025-35/SUBIT/SHM

Informations provided by the client

Sample Name: CBD OIL 20%
Matrix: Oil
Product: Oil

Laboratory Information

Acceptance Date: 03/09/2025
Test Start Date: 15/09/2025
Sampling: Sample delivered by the customer
Sample ID: 0030925149/SUBIT

Prepared for

CANNA HEALTH AMSTERDAM
28b Nieuwe Nieuwstraat Amsterdam
VAT NL002505280B09

Compound	Result	UM	LOQ	Measurement uncertainty
Arsenic (As) MS ^	< LOQ	mg/kg	0,05	
Cadmium (Cd) MS ^	< LOQ	mg/kg	0,01	
Mercury (Hg) MS ^	< LOQ	mg/kg	0,005	
Lead (Pb) MS ^	< LOQ	mg/kg	0,01	

%(m/m) = (Mass of the analyte/Mass of the product as it is)
NR Not Detected
LOD Limit of detection
LOQ Limit of quantification
<LOQ Below the limit of quantification

^Subcontracted
Method: MP/C/57 rev 4 2023, UNI EN 13805:2014 + UNI EN 15763:2010, MP/C/13 rev 3 2023

149/SUBIT



End of Test Report # Certificate of Analysis # 2025-35/SUBIT/SHM

Test end date: 16/09/2025
Issuing date: 30/09/2025

The results reported in the Test Report refer only to the sample actually tested, as received. The Laboratory declines all responsibility for the methods of carrying out sampling, transport and storage of the samples until delivery, if carried out by the customer. The Laboratory assumes responsibility for the information and data contained in the test report, with the exclusion of what is declared by the customer. This CoA can only be reproduced in complete form: if not complete, it is allowed only after written authorization from the Laboratory Manager. Copies of this Test Report and related documents are kept for 4 years. The laboratory declines responsibility for any information provided by the customer that may affect the validity of the results. Test Report digitally signed and compliant pursuant to art. 23 Legislative Decree 7 March 2005 No. 82 CAD and subsequent amendments and additions.



Certificate of Analysis # 2025-40/SUBIT/SMICRO

Informations provided by the client

Sample Name: CBD OIL 20%
Matrix: Oil
Product: Oil

Laboratory Information

Acceptance Date: 03/09/2025
Test Start Date: 11/09/2025
Sampling: Sample delivered by the customer
Sample ID: 0030925150/SUBIT

Prepared for

CANNA HEALTH AMSTERDAM
28b Nieuwe Nieuwstraat Amsterdam
VAT NL002505280B09

Compound	Result	UM	LOQ	Measurement uncertainty
Count of Microorganism at 30°C ^	< LOQ	UFC/g	10	
Count of Staphylococcus Aureus and other species coagulase positive ^	< LOQ	UFC/g	20	
Count of Escherichia coli beta glucuronidasi positive ^	< LOQ	UFC/g	10	
Count of Pseudomonas aeruginosa ^	< LOQ	UFC/g	20	
Mold count ^	< LOQ	UFC/g	20	
Yeast count ^	< LOQ	UFC/g	20	
Count of Enterobacteriaceae ^	< LOQ	UFC/g	10	
Salmonella ^	NR	Pres/Abs in 25g		

%(m/m) = (Mass of the analyte/Mass of the product as it is)
NR Not Detected
LOD Limit of detection
LOQ Limit of quantification
<LOQ Below the limit of quantification

^ Subappalto
Metodi: UNI EN ISO 4833-1:201; ISO 4832:2006; ISO 21528-2:2017; ISO 16649-2:2001; UNI EN ISO 6888-2:2021; UNI EN ISO 6579-1:2017; ISO 21527-2:2008

150/SUBIT



End of Test Report # Certificate of Analysis # 2025-40/SUBIT/SMICRO

Test end date: 12/09/2025
Issuing date: 30/09/2025

The results reported in the Test Report refer only to the sample actually tested, as received. The Laboratory declines all responsibility for the methods of carrying out sampling, transport and storage of the samples until delivery, if carried out by the customer. The Laboratory assumes responsibility for the information and data contained in the test report, with the exclusion of what is declared by the customer. This CoA can only be reproduced in complete form: if not complete, it is allowed only after written authorization from the Laboratory Manager. Copies of this Test Report and related documents are kept for 4 years. The laboratory declines responsibility for any information provided by the customer that may affect the validity of the results. Test Report digitally signed and compliant pursuant to art. 23 Legislative Decree 7 March 2005 No. 82 CAD and subsequent amendments and additions.