

HEALTH, NUTRITION & LIFESTYLE

GENERAL LIABILITY AND PRODUCT LIABILITY APPLICATION

		APPLICANT INFOR	RMATION
Applicant N	Name:		
Mailing Ad	dress:		
	City:	State:	Zip Code:
Location Ad	dress:		
	City:	State:	Zip Code:
We	ebsite:		Proposed Effective Date:
			From: To:
			12:01 AM Standard Time at the address of the Applica
pplicant is:	☐ Individual	☐ Joint Venture	LLC
	☐ Corporation	Partnership	Other - Specify:
	ness under current and puisitions of companies and	prior names I operations in the past 5 years	5):
		YOUR OPERAT	IONS
			10113
) Description o	of operations/list produc	ts and goods:	

3)	refrentage of your gross sales generated by the following types of operations		
a.	Manufacturer		%
b.	Contract-Manufacturer - Products sold under label of others		%
c.	Wholesaler/Distributor – Products of others sold under label of others		%
d.	Importer (Note: Products shipped directly to your customers without physical possession will not be considered as an acceptable form of business.)		%
e.	Retailer – Own label		%
f.	Retailer – Products of others sold under label of others		%
g.	Direct to customers via internet		%
h.	Other (please describe):		%
4)	If you are a Manufacturer, Contract-Manufacturer or Retailer – Own Label:		
a.	Have you or will you use ingredients imported from foreign suppliers?	Yes	☐ No
b.	Do you contract the manufacturing of your product to others?	Yes	No
If :	yes, please provide the manufacturer's name and physical address:		
5)	If you are a Wholesaler/Distributor - Products of Others Sold Under Labels of Others:		
a.	Please list the manufacturers and their physical addresses:		
b.	Do your suppliers each provide you with a certificate of liability insurance?	Yes	□No
c.	Do your suppliers also each provide you with additional insured-vendors coverage?	Yes	□No
6)	If you are an Importer, please list the countries of origin:		
7)	If you are a Contract-Manufacturer – Products Sold Under Label of Others:		
a.	What is the percentage of such products that are formulated entirely by the customer?		%
b.	Percentage of overall sales that consist of products sold under the labels of your customers?		%
c.	Do you have a written contract with each customer that includes hold harmless and indemnification agreements in your favor?	Yes	No
d.	Do you exclusively use ingredients supplied by your customer?	Yes	No

8)	If you	are a	Contract	-Packager	- For	Others:
----	--------	-------	----------	-----------	-------	---------

	YOUR PRO	DUCT SALES			
Annual Gross Sales:	Total	United States		Foreign	
Upcoming Year					
Current Year					
First Prior Year					
Second Prior Year					
Third Prior Year					
			Upcoming Year (Estimate):	Prior Yea tual	-
For use by children			-	%	(
Caffeine exceeding 300 mg per ser	ving (all sources)		-	%	(
Animal & vet supplements				%	
Sports nutrition - bodybuilding, m	uscle enhancement supple	ements		%	
Weight loss supplements			-	%	
exual enhancement supplements				%	(
Cannabidiol (CBD)/hemp products	8			%	(
TE: Coverage will not apply to procluding any product, supplement, ntrol Act of 1990 including amend 3 - Dimethylamylamine); Ephedra	additive, substance, ingred ments thereto, or the Anab Ephedrine Alkaloids; or Fe	ient or compound controlle olic Steroid Control Act of 2 nfluramine (N-Nitroso-Fenfl	d or banned by th 005; DMAA (Dimet uramine).	e Anabolic Si hylamylamin	teroid
		ND REGULATORY	COMPLIAN	ICE	
) Product Withdrawal/Product I					
Do you have a formal written pro	duct recall procedure?			Yes	
Have you voluntarily or involunta	rily recalled or withdrawn,	or are you considering reca	lling or	Yes	
withdrawing any products for an	y reason?				

11	L) Current practices or your specified industry equivalent:		
a.	Are you fully compliant with FDA Current Good Manufacturing Practices (cGMP)?	Yes	□No
b.	Are you compliant with Food, Drug & Cosmetic Act 21 CFR 111?	Yes	□No
12	Quality Assurance Program (QAP)/Quality Control Program (QCP):		
a.	Have you attained ISO 9000, QS 9000 or similar third party certification for your quality systems?	Yes	□No
b.	Do you have a formal written QAP/QCP, including written SOP's that control your operations?	Yes	□No
c.	Please provide name, title and contact information (email/phone) for QAP/QCP manager:		
13	3) Are all facilities used to manufacture, process, pack, hold or store your products registered with the FDA?	Yes	☐ No
14	4) If you are making or selling any Cannabidiol (CBD) products, are they tested and certified by a third party laboratory?	Yes	□No
a.	Do you have batch records on file that document production details for each lot of finished product?	Yes	No
b.	Are your products certified to contain no more than 0.3% THC and is it listed on the label?	Yes	□No
15	5) Labels:		
а.	Has outside legal counsel reviewed your labeling and confirmed it is in compliance with the regulations established by the FDA and FTC?	Yes	No
b.	Do all of your labels include a disclaimer that the FDA has not evaluated the claims on your labels and that your products are not intended to diagnose, treat, cure or prevent any disease?	Yes	□No
c.	Are you making any structure/function claims for your products on labels, websites or other marketing materials?	Yes	No
d.	Do you maintain documentation that substantiates each claim you make?	Yes	No
e.	Have you conducted, or are you planning to conduct, human clinical trials to substantiate your product claims?	Yes	No
	REGULATORY EVENTS		
16	5) In the past five years, have you submitted a Serious Adverse Event Report (SAER) to the FDA or has the FDA notified you of an SAER submitted directly by a health care provider, firm or consumer?	Yes	□ No
If	yes, please attach a comprehensive list of all Serious Adverse Events, along with copies of all reports and releve	ınt documei ———	nts.
17	7) Do you have an SOP detailing how to identify and handle an SAER/SAE?	Yes	□No

18) Are you aware of any complaint or notice filed in the last three years with any governmental agency or industry regulatory body, including but not limited to the FDA or FTC, concerning your product?	Yes	□No
If yes, please provide a detailed explanation:		
19) Have you been inspected by the FDA?	Yes	No
a. Did the FDA issue a Form 483 Notice of Inspection?	Yes	□No
If yes, please provide a copy.		
b. Are you aware of any study, analysis or trial conducted by the FDA or any industry regulatory body, to examine the safety of your products?	Yes	□No
c. Has FDA Form 483 been responded to with an FDA closeout letter?	Yes	No
20) Do you comply with Prop 65 labeling requirements?	Yes	□No
OPTIONAL COVERAGE ENHANCEMENTS		
21) Hired & Non-Owned Auto		
Check this box if Applicant would like to be considered for Hired & Non-Owned Auto Liability (HNOA) cover	age	
No, Applicant has separate Auto Policy		
□ N/A		
a. Do you own any auto that is used in your business and is registered to your Company?	Yes	□No
If yes, HNOA coverage is unavailable		
b. Does the Applicant have more than five employees using their personal auto for business use?	Yes	□No
If yes, HNOA coverage is unavailable		
c. Will any vehicle be operated beyond a 50 mile radius of the business location address on a weekly basis?	Yes	No
d. Will any vehicle be used for product delivery?	Yes	□No
If yes, HNOA coverage is unavailable		
22) Employee Benefits (Retro Date will be inception of our policy unless an expiring policy Declarations page is provided to document an earlier date)	Yes	□No

YOUR CLAIMS, LOSSES, DEMANDS FOR DAMAGES AND SIMILAR EXPERIENCE

Check here if no insure	d losses in the past 5	years				
defect in any prod	duct or work, which		suit, legal action or susp sult in a demand for dam s history?		Yes	□ No
If yes, please provide a	detailed explanation	:				
24) Current Carrier:						
Is current carrier offeri	ng renewal?				Yes	□No
Coverage Form:	Occurrence	Claims-Made	If Claims-Made, Retr	oactive Date:		
Limits:	\$		Deductible:	\$		
Premium:	\$		Rate:	\$		
25) Desired Limits:	ć		Desired Deduc	Aible. È		



Please initial:	
	en suppressed or misstated. I/We understand that this is an emission of this Application does not bind the Company to sell nor knowledge that any contract of insurance issued by the Company
Any person who knowingly and with intent to defraud any insu or statement of claim containing any materially false informatic concerning any material fact, commits a fraudulent insurance a	
Please initial:	
I/We hereby declare that the above statements and particulars any contract of insurance issued by the Company in response to	are true and I/We agree that this Application shall be the basis for o it.
Electronic signature of Applicant or Authorized Represer	ntative:
Title:	Current Date:
If you prefer not to return application with an electronic	signature, please print and sign below.
Signature of Applicant or Authorized Representative:	
Title:	Current Date:

Certaii	n terms are abbreviated in this application. Here are a few:
	FDA means the United States Food and Drug Administration
	FDCA-21CFR Part 11 means Food Drug and Cosmetic Act
	FTC means the United States Federal Trade Commission
	QAP / QCP means Quality Assurance Program / Quality Control Program
	SOP means Standard Operating Procedure
	cGMP / GMP means Current Good Manufacturing Practices / Good Manufacturing Practices
	Cannabidiol (CBD) is a psychoactive ingredient found in plant species cannabis sativa Prop 65 refers to the
	Safe Drinking Water and Toxic Enforcement Act of 1986
	tailed information on regulatory requirements and definitions, you may find useful references at www.fda d www.ftc.gov.
gov an	d www.ftc.gov.
gov an	
gov an	d www.ftc.gov.
gov an	d www.ftc.gov.
gov an	d www.ftc.gov.