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## APPLICATION FOR CLINICAL RESEARCH ORGANIZATIONS & CLINICAL TRIALS FOR PROFESSIONAL AND GENERAL LIABILITY INCLUDING PRODUCTS LIABILITY INSURANCE (Claims Made Basis)

## **APPLICANT'S INSTRUCTIONS:**

- 1. Answer all questions. If the answer requires detail, please attach a separate sheet.
  - 2. Application must be signed and dated by owner, partner or officer.
- 3. Please do not complete application earlier than 45 days before proposed effective date of coverage.
  - 4. PLEASE READ CAREFULLY THE STATEMENTS AT THE END OF THIS APPLICATION.
    (PLEASE TYPE OR PRINT IN INK)

		(I EDIOC THE ONT MINING)								
1.	APF	ICANT INFORMATION								
	a.	Full name of Applicant:								
	b.	Principal business premise address:								
		(Street) (County)								
		(City) (State) (Zip)								
	C.	Number of Employees: Full time Part time Seasonal Total								
	d.	Additional office locations:								
	e.	Name of parent company:								
	f.	Please describe all operations to be insured:								
	g.	Phone: ( )								
	h.	[ ] Corporation [ ] Partnership [ ] Joint Venture [ ] Sole Proprietor [ ] Other								
	i.	Date Established:								
2.	APF	ICANT OPERATIONS								
	a.	Fees and Receipts								
		Estimate for Estimate for Next								
		Current Year         Fiscal Year           Date: Fromto								
	b.	Percentage of foreign professional services and provide the names of the countries involved:								
	C.	Do you manufacture or sell any products?								
	C.	If Yes, please attach a detailed description of your current products and any future products being researched.								
	d.									
		(i) Please describe this phase:								
		(ii) Will this phase be performed in accordance with an FDA approved protocol?								
		(iii) Please indicate IND number:								
		(iv) Will this phase and have all previous related phases been performed in accordance with an FDA approved protocol?								

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	e.	Will you or your employees provide any health care services in conjunction with this trial? [ ] Yes [ ] No If Yes: Professional Title:							
		Description of services provided:							
,	f.	Is the clinical investigator an employee of your firm?							
	g.	Is the clinical investigator an employee of the test site facility?							
	h.	(i) Please provide the name and the proposed use or function of the product being tested.							
		(ii) Are you aware of any other approved uses or functions of the product being tested?							
		(iii) Do you have any knowledge that this product or any of its components might cause or contribute to any immune system reactions?							
i	i.	Please provide the name of the product manufacturer (if other than yourself):							
j	j.	Is the Applicant a "Covered Entity" under the Health Insurance Portability and Accountability Act of 1996 (HIPAA) Privacy Rule?							
		If Yes,							
		(i) Has the Applicant implemented procedures to comply with the HIPAA Privacy Rule? [ ] Yes [ ] No							
		(ii) Provide the name and title of the Applicant's Privacy Officer.							
		Our Business Associate Agreement is available at <a href="https://www.markelcorp.com/PolicyholderServices">www.markelcorp.com/PolicyholderServices</a> . This is the only Business Associate Agreement we will recognize.							
3.	TES	STING INFORMATION							
;	a.	Please indicate the anticipated number of test subjects over the next 12 months:							
	b.	Please give the sex and age of the test subjects:							
	C.	How will test subjects be recruited? Please provide a detailed explanation.							
	d.	Will test subjects be required to sign an informed consent document?							
	а. e.	, , , , , , , , , , , , , , , , , , , ,							
	f.	The anticipated trial period: From To To How will the trial be conducted and by whom?							
,	g.	Please attach a detailed explanation.  How will the trial be funded?							
	h								
	h.	Where will the trial be performed? Please check the appropriate response.  [ ] Facility & Location [ ] Non-Profit Testing Institute [ ] Clinical Research Center [ ] Other (please describe)							
	i.	(i) Will an Institutional Review Board oversee the trials?							
		(ii) Are you a member of this Board?							
j	j.	Please indicate the number of employed professionals or independent contractors. (IF NONE, STATE NONE.)							
		Contractor <u>Employee</u> <u>Independent</u> <u>Total</u> (i) RN/LPN							
		(i) RN/LPN							
		(iii) Clinical Investigator							
		(iv) Clinical Research Assoc.							

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		<u>Employee</u>	Contra <u>Indeper</u>		<u>Total</u>	
	(v) Physician	<u>=111610 y 00</u>	шаорог	<del>Idorit</del>	<u>rotar</u>	
	(vi) Medical Monitor					
	(vii) Engineer			<del></del>		
	(viii) Biostatistician					
	(ix) Data Entry			<u> </u>		
	(x) Legal Counsel					
	(xi) Other					
k.	Do you perform any environn If Yes, please attach a detaile			ng?		[ ]Yes [ ]N
l.	Please indicate testing perfor over the next 12 months:	med on specif	ied product	s over the last	12 months and	anticipated testing to be performe
		La		Next		
	(1)	12 Mc	intns	12 Months		
	(i) Hormones & Steroids					
	(ii) Vaccines					
	(iii) Injectables					
	(iv) Prescription Products	1				
	(v) Over the Counter					
	(vi) Diet Aids					
	(vii) Vitamins					
	(viii) Food Supplements					
	(ix) Novel Drugs					
	(x) Generic Off-Patient					
	(xi) Products, Other than A	bove				
	(xii) Instruments (x-diagnos	tic)				
	(xiii) Cosmetics, Health & Beauty Aids					
	(xiv) Surgical Equipment					
	(xv) Diagnostic Instruments & Equipment					
	(xvi) Therapeutic Devices					
	(xvii) Life Support					
	(xviii) Other					
ΔPF	PLICANT HISTORY					
a.	Provide a brief description of	the results of	any previo	us related trial	s:	
<b></b>						
b.	Fully describe any adverse re	esults from pre	evious relat	ed trials includ	ling animal stud	lies and/or toxicity studies:
C.	List any claims related inform	ation provide	d in 4(a) an	d 4(b) above:		
	Dat			Land 9	N	6 Indiana
	<u>Claimant</u> <u>of Lo</u>	<u>ss</u> <u>Ex</u>	<u>pense</u>	Indemnity	<u>Nature o</u>	<u>r injury</u>

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5.	CLAIMS  (Attach a detailed explanation for any "Yes" answers)									
	a.									]Yes [ ]No
	b.							d & Drug Administrati ogics Evaluation and		]Yes [ ]No
	C.							ny federal, state or loc		]Yes [ ]No
	d. Do you operate in compliance with the FDA's Good Clinical Practice Guidelines?							[	] Yes [ ] No	
	e.							cal Practices or any fe		]Yes [ ]No
6.	COV	/ERA	GE							
	a.	Lim	its of liability desire	d: \$						
	b.	Amo	ount of deductible d	esired:	\$		_			
	C.	Pres	sent coverage							
		<u>Car</u>	<u>rier</u>	<u>Prof</u>	<u>GL</u>	Deductibl	e/SIR	Limits	Claims I Yes	Made? No
		_								
		If Ye	es, please provide a	an expla	nation.					
	d.	Ret	roactive date (if app	licable)						
7.	ADE	OITIO	NAL INFORMATIO	N						
	Please provide the following information with this application:									
		(i)	Advertisements, b							
		<ul> <li>(ii) Sample contract between you and the clinical trial investigator, if the investigator is not your employee of the test site facility.</li> </ul>								mployee or a
		(iii)	Informed consent	docume	ent.					
		(iv)	Most recent Annu	al Repo	rt or audite	ed financial s	atement			
	(v) Copy of letterhead or other business stationary.									
"CL	AIMS I	MADE	" basis for ONLY	THOSE	CLAIMS T	THAT ARE F	IRST MAD	TED IN THE POLICY DE AGAINST THE IN ance with the terms of	SURED DURING	
here	in is tr	ue an	d that it shall be the	basis of	the policy	of insurance	and deem	notice stated above ar ed incorporated therei	n, should the Insu	rer evidence it
			is application by iss ig manager, Comp				ize the rel	lease of claim inform	nation from any p	rior insurer to
Nan	ne of A	pplica	ant*				Title (Offi	cer, partner, etc.)		
Sigr	Signature of Applicant*					Date				

Signing this application does not bind the Applicant or the Insurer or the Underwriting Manager to complete the insurance, but one copy of this application will be attached to the policy, if issued.

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