MEDICAL DEVICE MANUFACTURING LICENSE APPLICATION PLEASE COMPLETE THIS FORM FULLY—INCOMPLETE APPLICATIONS WILL BE RETURNED See Page 3 for Instructions

License Number (if not new):

			SHIP CHANG		Address:	, , , , , , , , , , , , , , , , , , ,		
	Name of Firm				Mailing Address (if dit	fferent or P.	O. Box n	umber)
2.	2. DBA (Use other sheets as needed)			7. Mailing Address (continued)				
3.	Facility Address (number, stre	eet)		8. City		State	ZIP Code	
4.	4. Facility Address (continued)			9. Country (if other than United States)				
5.	City	State	ZIP Code	10.	Website (URL)			
	. Interstate Commerce Product Shipped . Type ownership * Please atta Individual/Sole Proprietorsh			rship	aterials Received * Corporation	Limited	N/A	Corporation
10	Non Profit	•	Other					
	. Owner's Name/Corporate Na				State of Incorporati			
14	. Owner's or Officers' Names a	nd Titles			Owner's or Officers	s' Names ar	nd Titles	
45	The state of the second			L .)			. ()	
15	. Type of Manufacturing Busine Manufacturer Con	ess (Cne itractor	ск ан tnat app Compone	• ·	Size of Facility Number of En			ity.
	Specification Developer Other:			110	Business Day			
16	. Stage of Manufacture at Date			all th				
	Manufacturing Design Products Develop		Design Validation		Pre-Production Design Transfer	Other:		
17.	-	tination (n Clinical igational	Trials Ca	aliforn		Distribution	E	xport Market
18	. Check Each Product Area tha	t Applies	s to devices M	lanufa	actured			
862 Clinical Chemistry and Toxicology 874			Ear,	Nose, and Throat		8	86 Ophthalmic	
	864 Hematology and Pathology 876			76 Gastroenterology/Urology 888		88 Orthopedic		
	866 Immunology and Microbiology 878			8 Gen	eral and Plastic Surg	ery	8	90 Physical Medicine
	868 Anesthesiology				eral Hospital and Per	rsonal Use	8	92 Radiology
	870 Cardiovascular				rological			
	872 Dental 884 O			Obst	Obstetrical and Gynecological			

PLEASE CONTINUE TO NEXT PAGE

MEDICAL DEVICE MANUFACTURING LICENSE APPLICATION

	License Number (if not new):							
		es of classif cessary.	ied and/or uncla	assified medi	cal devices manufactured in th			
Federal Classification and Title CLASSIFICATION (Ch							heck One)	
reacta		Sincution					II	
					manufacture of the devices lis	ted above a	and if the activ	vities will
					eets if necessary.			
		Activities	In-House	Contract	Process or Activities	S	In-House	Contract
Sterilizati					Repackaging/Relabeling			
Software		•			Remanufacturing/Refurbishin	ng		
Circuit Bo		ssembly			Tissue/Cell Culture			
Lyophiliza					Other:			
Antigen/A	Antiboo	dies						
21. Paym	nent Co	ode (Check	all that apply)					
А	\$4,7	\$4,187.00 Base Fee Fee Fee is due at the time application is submitted and is NON- REFUNDABLE				NON-		
В	\$	10.00	00 Late Fee If over 30 days past license expiration date					
\$ TOTAL AMOUNT DUE Payable to: CA Department of Public Health								
22. If you	ı meet	one of the l	isted criteria be	low FDB can	issue a license without first ir	nspecting ye	our facility.	
Check	the a	ppropriate b	box(es) and atta	ach documen	tary evidence.			
Ac	opy of	f a valid biol	ogics license is	sued by the l	J.S. Food and Drug Administr	ation (FDA)	
A copy of a valid establishment registration pursuant to Section 510 of the federal act and attestation that a								
federal inspection was completed within the last two years								
A copy of documentation demonstrating compliance with audits conducted pursuant to International Organization								
for Standardization (ISO) ISO standards (ISO 9000 series, ISO 13485:2003, ISO 15378:2006)								
A copy of approved investigational device exemption issued by the FDA								

None of the above apply (Inspection will be required)

The Food and Drug Branch **MUST BE NOTIFIED IMMEDIATELY** of any changes in the above information as provided by California Health and Safety Code, Section 111630. Under penalties of perjury, I declare that the information included with this application and all attachments are true, correct, and complete. I also give permission for the below authorized representatives and/or signatories to speak about the application with CDPH.

23.Owner's Signature	Owner's Printed Name	Title Owner/	Date
Auth	orized Representativ	es and/or Signatories	S:
04 Duraimana Omeratan Manaa	OF Talaukawa Nuushaw		

24. Business Operator Name	25. Telephone Number	26. Emergency Number	27. E-Mail Address
28. Correspondent Name	29. Telephone Number	30. Alt Phone Number	31. E-Mail Address

End of Application. Please note: All boxes must be completed.

License Number	Expiration Date	Date Received	Payment Type	Amount \$
				Ŷ

MEDICAL MANUFACTURING LICENSE APPLICATION INSTRUCTIONS

A separate application is required for each place of business. Please complete and/or amend this application as is most appropriate to your facility. Include the appropriate fee for each application and make payable to: <u>CA DEPARTMENT OF PUBLIC HEALTH</u>. The fee must accompany this application or it cannot be processed. Unsigned or incomplete applications cannot be processed. The following are further instructions on how to complete this application:

New Applicant: Place an (X) in the box next to New Applicant if your firm has not previously applied for a Medical Device Manufacturing License at this location while under the current ownership. **This license is non-transferable.** If your firm has changed location, ownership, or both, place an (X) in the appropriate box **and also** in the box next to New Applicant. For renewal applications, place an (X) in the box next to Renewal. For any section that does not apply to your company, please indicate with (N/A). **Do not leave any sections blank**.

- 1. **Name of Firm:** Enter full name of business, corporation, company, or organization applying for licensure.
- 2. **DBA:** Enter any other name(s) your company is doing business as.
- 3.–5. Facility Address: Enter the number, street, city, state, and ZIP code for this facility location.
- 6.–8. **Mailing Address:** Enter the full mailing address if different from the facility address or P.O.Box.
 - 9. **Country:** Enter the country where your facility is located.
- 10. Website: Enter the website address for your business, if applicable.
- 11. **Interstate Commerce:** Place an (X) in all appropriate boxes that correctly describe your business' receipt or distribution of products or materials through or into interstate commerce.
- 12. **Type of ownership:** Place an (X) in the box next to the appropriate legal description of the facility's ownership. Attach evidence of ownership.
- 13. **Owner's or Corporate Name. State of Incorporation:** List the business owner's name or corporate name. Identify the state of incorporation.
- 14. **Owner's or Officers' Names and Titles:** List other business owner's names or officers' names and titles. Use additional sheets if necessary.
- 15. **Type of Manufacturing Business:** Place an (X) in the box next to each type of manufacturing business conducted at this facility, size of facility, number of employees, and list business days and hours.
- 16. **Stage of Manufacture:** Place an (X) in the box next to the stage of manufacture your products are in at the time of application submission. Check all that apply.
- 17. **Intended Device Destination:** Place an (X) in the box adjacent to the destination(s) for your manufactured products. Check all that apply.
- 18. **Products Manufactured:** Place an (X) in the box adjacent to each product that applies to the devices manufactured or to be manufactured.
- 19. **Classified or Unclassified Products Manufactured:** For each medical device product, list the federal classification name and classification category (I, II, or III) as listed in 21 CFR, Sections 862 to 892.
- 20. **Manufacturing Processes:** Place an (X) in the column adjacent to any indicated processes to identify if they will be done in-house or contracted out. Leave line blank if the indicated process will not be used in the manufacture of listed devices. List additional processes or methods as needed herein or on additional sheets, if necessary.

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MEDICAL MANUFACTURING LICENSE APPLICATION INSTRUCTIONS

21. **Payment Code:** Place an (X) in the box adjacent to each applicable fee.

Payment Type	Fee	Description	Payment Code
New, Relocation, Ownership Change, or Renewal	\$4,187	Base fee, due at time of application	А
Late Fee	\$10	Due if over 30 days past license expiration date	В

LICENSE FEES ARE NON-REFUNDABLE AND NON-TRANSFERABLE TO OTHER LOCATIONS OR ENTITIES

- a. Enter license fee according to payment codes above. (License valid for 2 years)
- **b.** Enter Total Payment Due by adding "A" and "B".
- 22. **Inspection Criteria:** Attach U.S. Food and Drug (FDA) or International Organization for Standardization (ISO) Standard Documents. Place an (X) in the appropriate box(es) for the items that you are submitting with the application. For more information regarding this requirement refer to the California Health and Safety Code Section 111635. If you meet any of the listed criteria on the application by providing documentary evidence, FDB may be able to issue a license without first inspecting your place of business. The documentary evidence must be valid for your current establishment. If no criteria are met, an inspection will be required.
- 23. **Owner's Signature:** Sign the application. Print the signer's signature, title, and indicate signature date. All signatures must be original.
- 24.-27.**Business Information:** Print the business operator's name, title, business and emergency telephone number, and E-Mail address.
- 28.-31.**Correspondent Information:** Please print the correspondent's name, title, telephone number, and email address.

	Please make all checks payable	to: CA Dep	artment of Public Health				
	Mail Application and checks to:						
Regular Mail:	California Department of Public Health	Overnight Mail:	California Department of Public Health				
	Food and Drug Branch – Cashier MS 7602 P.O. Box 997435 Sacramento, CA 95899-7435		Food and Drug Branch – Cashier 1500 Capitol Avenue, MS-7602 Sacramento, CA 95814				

If you have any questions about this application, please contact the FDB License Desk for Drug Manufacturing at (916) 650-6500, (800) 495-3232.