In accordance with the California Health and Safety Code, Section 119313, a body art facility shall maintain and follow a written Infection Prevention and Control Plan, provided by the owner or established by the practitioners, specifying procedures to achieve compliance with the Safe Body Art Act. A Copy of the Infection Prevention and Control Plan shall be filed with the Local Enforcement Agency and a copy maintained in the body art facility.

The body art facility owner shall provide onsite training on the facility's Infection Prevention and Control Plan to the body art practitioners and employees or individuals involved with decontamination and sterilization procedures.

Training shall be provided when tasks where occupational exposures may occur are initially assigned, anytime there are changes in the procedures or tasks and when new technology is adopted for use in the body art facility, but not less than once each year. Records of training shall be maintained on-site for three years.

The Infection Prevention and Control Plan shall be maintained current and updated whenever there are changes to any of the procedures or tasks listed and when new technology is adopted for use in the facility.

Name of Body Art Facility:	
Site Address:	
City, State, ZIP:	
Type of Body Art Facility:	
Contact Person:	Telephone:

. D w	econtamination and Disinfection: Describe the procedures for decontaminating and disinfecting of orkstation and surfaces(California Health and Safety Code 119308 (b) and 119309 (a)(b)(c)(d)(e)).
1.	Workstation surfaces/counter tops:
2.	Workstation chairs/stools:
3.	Trays:
4.	Armrests:
5.	Headrests:
6.	Procedure area:
7.	Tables:

	8.	Tattoo machine and Clip Cord:						
	9.	Reusable instruments, calipers, needle tubes, etc. portable light fixtures or other:						
	10.	Permanent Cosmetic Machine:						
;	ste ste dis wh Co	eusable Instruments or Disposable: Describe the procedures used for decontaminating, erilizing, packaging and storing of reusable instruments. Include the procedures for labeling of erilized peel-pack. Indicate whether the body art facility uses all pre-sterilized, single-use and sposable instruments. Describe the record keeping logs and procedure logs maintained on-site nen using 100% pre-sterilized, single-use and disposable instruments (California Health and Safety ode 119309 and 119315). Needle tubes:						
	2.	Calipers:						
	3.	Other instruments:						

C.	ins	Storage: Describe the storage location and equipment used for the storage of clean and sterilized instrument peel packs to protect the packages from exposure to dust and moisture (California Health and Safety Code 119315 (c)).						
	th	et Up and Tear Down of Workstation: Describe the procedure for setting up and tearing down e workstation for the following procedures (California Health and Safety Code 119308, 119309), 119311, and 119313 (b)(4)).						
	1.	Tattoo:						
	2.	Piercing:						
	3.	Permanent Cosmetics:						
	4.	Branding:						
E.	of inl du pr	revention of Cross Contamination: Describe the techniques used to prevent the contamination instruments, tattoo machines, trays, tables, chairs, clip cords, power supplies, squeeze bottles, ks, pigments, lamps, stools, soaps, procedure sitesand additional areas of potential contamination uring body art procedures. Include barriers provided to prevent cross contamination. Describe how ocedure sitesare prepared for a body art procedures. (California health and Safety Code 119308, 9309, and 119311 (c)(d)(e)(f)).						

E.	Prevention of Cross Contamination (Continued):								
F.	lo	narps Containers: Describe the procedures used for the safe handling of sharps and indicate the cation of the in-use sharps containers. Indicate disposal frequency for sharps waste (California ealth and Safety Code 119314 (e)).							
G.		narps Disposal: Describe the disposal of sharps used during a body art procedure (California ealth and Safety Code 119308 (b)(3) and 119311 (g)).							
	1.	Needles and needle bars:							
	2.	Razors:							
	3.	Other sharps or single-use marking pens used on open skin:							
Н.		st the Medical Waste Hauler, Mail-back System or Alternative Treatment Technology used r the disposal of sharps containers (California Health and Safety Code 119314 (e)):							
		Andinal Wanta Haular							
		Medical Waste Hauler:							
		Sity State 7IP:							

afety Code 119301 (k) and 119308 (b)(6)).
and Temperature: List the temperature of the autoclave and duration of time at that rature required for the sterilization of clean instruments. Indicate wherethe sterilization log is ained on-site. Indicate whether each sterilization load is tested using Class 5 integrators ornia health and Safety Code 119315 (b)(3)(5)).
erature:
nal Protective Equipment: List the personal protective equipment used during a body and dure for the practitioner and the client (California Health and Safety Code 119308 (a) and 9 (j)).
washing Sink: List the locations of the handwash sinks and describe the items supplied a sink (California Health and Safety Code 119314 (b)(3)).
k

Ο.	Aftercare Procedure: Describe the written recommendation and care information provided to the client after a body art procedure. List the type of bandages or wrapping provided after a body art procedure (California Health and Safety Code 119309 (a)(b)(c)).
P.	Procedure for an Accidental Spill: Describe the clean-up and disinfection procedure taken when there is an accidental spill of sharps (California Health and Safety Code 119309 (a)(b)(c)).
Q.	Trash Receptacles and Disposal of Contaminated Trash: List the type of trash receptacles used and their location throughout the body art facility. Describe the procedure for the disposal of contaminated items, such as gloves (California Health and Safety Code 119311 (a) and 119314 (d)).
R.	Negative/Failed Spore Test: Describe the procedure conducted when a monthly spore test has failed. Indicate where the facility maintains a spore test log on-site (California Health and Safety Code 119315 (b)(2)(4)).
S.	Commercial Ink or Pigment Manufacturers: List the manufacturer(s) for the inks or pigments used at the facility. Describe the procedure for dilution of inks. Only sterile water should be used for dilution of inks orpigments (California Health and Safety Code 119311 (b)(c)(d)(e)).

Т.	Permanent Cosmetic Machine Name and Manufacturer: Provide the model name and numbe for the permanent cosmetic machine(s) used (California Health and Safety Code 119311 (i)(j)).							
U.	Service Animals: Describe the facility's policy regarding service animal presence in procedure, decontamination, and sterilization areas (California Health and Safety Code 119314 (f)).							
	intain a copy of this completed document in your files. Submit one copy to the Local Enforcement ency.							
ind the	ereby certify that all body art practitioners performing bodyart at this facility and employees or dividuals involved with decontamination and sterilization procedures have been trained with e procedures and information contained in this document. To the best of my knowledge and lief, the statements made herein are correct and true.							
Sig	gnature: Date:							
Tit	le:							

Sterilization Procedures

When a body art facility is equipped with a decontamination and sterilization room and will be sterilizing reusable instruments and body art jewelry, the following sterilization procedures must be followed:

- Clean instruments to be sterilized shall first be sealed in peel-packs that contain either a sterilizer indicator or internal temperature indicator. The outside of the pack shall be labeled
- 2. Sterilizers shall be loaded, operated, decontaminated and maintained according to manufacturer' directions, and shall meet all of the following standards:
 - Only equipment manufactured for the sterilization of medical instruments shall be used.
 - Sterilization equipment shall be tested using a commercial biological indicator monitoring system after the initial installation, after any major repair, and at least once per month. The expiration date of the monitor shall be checked prior to each use.
 - Each sterilization load shall be monitored with mechanical indicators for time, temperature, pressure, and at a minimum, class V integrators. The Class V integrator gives an immediate response on whether the sterilization has been achieved. Each individual sterilization pack shall have an indicator.
 - Biological indicator monitoring test results shall be recorded in a log that shall be kept on site for two years after the date of the results.
 - A written log of each sterilization cycle shall be retained on site for two years and shall include all of the following information:
 - The date of the load.
 - A list of the contents of the load.
 - The exposure time and temperature.
 - The results of the Class V integrator.
 - For cycles where the results of the biological indicator monitoring test are positive, indicate how the items were cleaned, and proof of a negative test before reuse.
- 3. Clean instruments and sterilized instrument packs shall be placed in clean, dry, labeled containers, or stored in a labeled cabinet that is protected from dust and moisture. Use clean gloves to handle sterilized packages to prevent cross contamination of the sterilized item when the package is opened for use.
- 4. Sterilized instruments shall be store in the intact peel-packs or in the sterilization equipment cartridge until time of use.
- 5. Sterile instrument packs shall be evaluated at the time of storage and before use. If the integrity of a pack is compromised, including, but not limited to, cases where the pack is torn, punctured, wet or displaying any evidence of moisture contamination, the pack shall be discarded or reprocessed before use.

Sterilization Procedures

- 6. A body art facility that does not afford access to a decontamination and sterilization area that meets the standards of subdivision (c) of Section 119314 of the California Health and Safety Code or that does not have sterilization equipment shall use only purchased disposable, single-use, pre-sterilized instruments. In place of the requirements for maintaining sterilization records, the following records shall be kept and maintained for a minimum of 90 days following the use of the instruments at the site of practice for the purpose of verifying the use of disposable, single-use, pre-sterilized instruments:
 - A record of purchase and use of all single-use instruments.
 - A log of all procedure, including the names of the practitioner and client and the date of the procedure.

Operating Conditions for Autoclave

Cleaning: Remove all material on the instruments during the cleaning process to ensure that the sterilization process is achieved. The cleaning process can be a manual cleaning or by use of an ultrasonic machine.

Packaging: Package the instruments with hinges in the open position to ensure that the ridges and crevices of the instruments are sterilized.

Loading: Load the autoclave with the packages upright on their sides. Peel packs should be on edge with the plastic side next to a paper side to allow for steam penetration. Do not overload the autoclave to allow proper flow of the steam to achieve sterilization.

Steam Sterilization: Temperature should be 121° C or 250° F: pressure should be 106kPa (15lbs/in2); 30 minutes for packaged items. At a higher temperature of 132° C or 279° F, pressure should be 30 lbs/in2; 15 minutes for packaged items.

Allow all items to dry before removing them from the autoclave. Use clean gloves to handle packaged items.

Pressure settings (kPa or lbs/in2) may vary slightly depending on the autoclave used. Follow manufacturer's recommendations for your autoclave.

Exposure time begins only after the autoclave has reached the target temperature.

Source: Adopted from Principles and Methods of Sterilization in Health Sciences. JJ Perkins. 1983

Sterilization Log

Date	Load #	Contents	Operator	Time	Temp	Psi	Temp indicator Results	Attach Integrator	Spore Test Results	Action Taken due to Failed Results