INFECTION CONTROL PLAN

Facility Name:	Effective Date:
List of Services Offered:	

PART 1 - LIST OF EQUIPMENT/INSTRUMENTS USED (Additional page in Appendix B)

Equipment/Instruments Used		Supplier	Product Verification
i.e. inks, pigments, needles, cartridges		Manufacturer/Model	(attach a copy of verification document)
	☐ Single use ☐ Reused ☐ Sterilized Onsite		□Sterilization certificate (if pre-sterilized) Method:
			☐Backflow prevention (needle cartridges)
	☐ Single use ☐ Reused ☐ Sterilized Onsite		☐Sterilization certificate (if pre-sterilized) Method:
			☐Backflow prevention (needle cartridges)
	☐ Single use ☐ Reused ☐ Sterilized Onsite		□Sterilization certificate (if pre-sterilized) Method:
			☐Backflow prevention (needle cartridges)
	☐ Single use ☐ Reused ☐ Sterilized Onsite		☐Sterilization certificate (if pre-sterilized) Method:
	_ stermzed orisite		☐Backflow prevention (needle cartridges)
	☐ Single use ☐ Reused ☐ Sterilized Onsite		☐Sterilization certificate (if pre-sterilized) Method:
	E Stermized Offsite		☐Backflow prevention (needle cartridges)
	☐ Single use ☐ Reused		☐Sterilization certificate (if pre-sterilized) Method:
	☐ Sterilized Onsite		☐Backflow prevention (needle cartridges)
	☐ Single use ☐ Reused		☐Sterilization certificate (if pre-sterilized) Method:
	☐ Sterilized Onsite		☐Backflow prevention (needle cartridges)

Part 2 – CLEANING, STERILIZATION AND DISINFECTION PROCEDURES

SECTION A – STERILIZATION PROCEDURE

Critical Items	Critical Items								
Describe how steril	ized equipr	ment/instru	ments will be						
stored before use									
Describe how you v	vill monitor	for evnired	d nra-sterilized						
single use instrume		TOT CAPITE	a pre stermzeu						
Describe how expir	•	mised pre-	sterilized						
instruments will be	handled								
Onsite Sterilization	Equipmen	t							
☐ Autoclave			Brand:	Model:	CSA Approved:	International Certification:			
☐ Chemiclave					☐ Yes ☐ No	☐ Yes ☐ No			
☐ Other									
Monitoring Require	ements if S	terilizing O	nsite 						
Temperature:	Time:		Pressure:	Spore Testing:	Spore Test Frequency:				
				☐ Yes ☐ No					
Class 5 Process Indi	cator:	Colour Ex	l pected:	Laboratory Analysis :	Laboratory Name:				
☐ Yes ☐ No			P 00000.	☐ Yes ☐ No					
Describe your procedures in the event your sterilizer has									
malfunctioned or sterilization package has been									
compromised									

SECTION B – CLEANING AND DISINFECTION PROCEDURES

Semi/Noncritical Items	Semi/Noncritical Items									
Describe how instrumer	nts are held prio	r to cleaning and								
disinfecting and where o										
Describe how instrumer										
what tools are used to a	ssist with cleani	ng								
Describe how cleaned a	nd disinfected in	struments are								
stored										
Duaduat Nama	DIN	A ative In and diam't	Disinfestant	Describe Dress draws Instruction Consentration and Coals Time						
Product Name	DIN	Active Ingredient/ Target Organism	Disinfectant Level	Describe Procedures Including Concentration and Soak Time						
		Taiget Oigainsiii	Level							
			□Low							
			□Intermediate							
			□High							
			-							
			□Low							
			□Intermediate							
			□High							
			□Low							
			□Intermediate							
			□High							
			-							
			□Low							
			□Intermediate							
			□High							

SECTION C - SINGLE USE PROCEDURES

Single Use Items	
Describe method of disposal	
Describe how items will be stored	
Describe how pre-sterilized instruments will be monitored for expiration	

SECTION D – OTHER PRODUCTS USED (i.e. Lotion, antiseptic, etc)

Product Name	DIN	Active Ingredient	Describe How and When it is Used	Storage

SECTION F – LAUNDERING PROCEDURES

List of Laundered Items	Where is it Laundered	Frequency	Washing/Drying Settings	Storage

Part 3 – OPERATIONAL PROCEDURES

SECTION A – OPERATIONAL PROCEDURES (Use a separate sheet for each service provided)

Set Up and Tear Down Procedures			
Service Provided:			
Describe your set up procedures		Describe your tear down procedur	es
Barrier Use			
List the type of barrier used	List equipment/	body part barrier is used on	How often is it changed

SECTION B – CLIENT CARE (Attach a copy of your consent and aftercare forms)

Section B Chile (Account Copy of Your Consent and arcercare	10111137						
Client Care Procedures							
Service Provided:							
Describe how the area of the body is prepared	Describe aftercare procedures						
Bandages/Covering							
List the type of bandages and coverings used	Manufacturer						

APPENDIX A – SUMMARY TABLE FOR INSTRUMENT/EQUIPMENT CLASSIFICATION AND DETERMINING PRODUCT DISINFECTION LEVEL

Table 1. Instrument/Equipment Classification and Product Disinfection Level

Classification	Level of Sterilization/Disinfection	Instrument/Equipment	Determining Product Disinfection Level				
Critical	Sterilization	Any instrument/equipment intended to puncture the skin or contact the puncture site or a sterile instrument before puncturing					
Semicritical	High-Level Disinfection	Any instrument/equipment intended to contact nonintact skin or a mucous membrane but not penetrate it	When choosing a high-level disinfectant, make sure the manufacturer's label has a: • DIN • Disinfectant claim • "TB" claim and specifically states "high-level disinfectant" or "chemical sterilant" or "sporicidal". If it does not state this, then it is not acceptable unless it meets all requirements in Table 14: Infectious Agents Killed by Disinfection of the Guidelines for Personal Services Establishment.				
Noncritical	Intermediate-Level Disinfection	Any instrument/equipment intended to contact intact skin but may accidentally contact nonintact skin or receive blood or body fluid splatter	When choosing an intermediate-level disinfectant, make sure the manufacturer's label has a: DIN Disinfectant claim "TB", "tuberculodical" or "mycobacterium" claim. These disinfectants are equipped for higher-risk surfaces and equipment such as those that may come into contact with nonintact skin. They are harder to find in retail stores, but readily available from cosmetic, dental or medical supply companies.				
	Low-Level Disinfection	Any instrument/equipment that does not directly touch the client or contacts only intact skin	When choosing a low-level disinfectant, make sure the manufacturer's label has a: • DIN • General disinfectant claim Low-level disinfectants are suitable for surfaces and equipment that would only, at most, come into contact with the client's intact skin. This level of disinfectant is easy to find in retail stores.				

APPENDIX B – TEMPLATE FOR EQUIPMENT/INSTRUMENT LIST

Equipment/Instruments Used		Supplier	Product Verification
i.e. inks, pigments, needles, cartridges	_	Manufacturer/Model	(attach a copy of verification document)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)
	☐ Single use		☐Sterilization certificate (if pre-sterilized)
	☐ Reused		Method:
	☐ Sterilized Onsite		
			☐Backflow prevention (needle cartridges)

APPENDIX C – SAMPLE STERILIZATION LOG SHEET

Personal Service Establishment Name and Address:			Month/Year									
Equipment Name and Model Number:				Serial Number:								
Date	Time			Temp.	Pres	sure	Temp. Sen		Operator	's	Comments	
(dd/mm/yy)	Start	End	Cycle Length	∘F or ∘C			Indicator: Change Of	Colour oserved: Y/N	Initials			
☐ Monthly spore strip tests submitted ☐ Date ((dd/mm/yy)		☐ Re	sults	Date (dd/mm	/yy) 		Results	Date (dd/mm/yy)		

Indicate any corrective action taken on reverse. Use one operation log per sterilizer within the personal service establishment. This record should be kept for five years.