

(702) 486-7044 • (800) DDS-EXAM • Fax (702) 486-7046

INFECTION CONTROL INSPECTION/SURVEY FORM					Rev 1	0/2014			
Dental Office Name:					Date of Inspection:				
Lice	nsee Name:		Owner Dentist:						
Address:				INSPECTOR(S)					
				(1) (2)					
City		State:	Zip Code:	PUR	POSE OF INSPECTION				
		Nevada		Initial Inspection:	Random	Inspec	tion:		
		COMP	LIANCE LEVEL C	RITERIA – LEVEL # 1-4					
#1-	# 1 - CRITICAL: MUST BE MET. COULD RESULT IN IMMEDIATE TERMINATION OF PATIENT CARE AND EXTENDED OFFICE INABILITY TO TREAT PATIENTS.								
#2	- REMEDIAL ACTION REQUIRED	REQUIRES COF		IANCE WITHIN 7DAYS.					
#3	- ACTION REQUIRED: REQUIR	ES CORRECTIVE		WITHIN 30 DAYS.					
# 4 - ACTION RECOMMENDED: NOT REQUIRED FOR COMPLIANCE AT THIS TIME – COMPLIANCE REQUIREMENTS SUBJECT TO CHANGE AS CENTER FOR DISEASE CONTROL (CDC) REQUIREMENTS MAY CHANGE.									
REC	CORD KEEPING – EACH PR					LEVEL 1-4	Y	Ν	
1	Written infection control progra	im that is specific	for the owner of	this location		3	Y	N	
EDU	ICATION & TRAINING								
2	Documentation of review of the	infection contro	l plan at least ann	ually to ensure compliance w	vith best practices	3	Y	Ν	
3	Documentation of Bloodborne F	Pathagons trainin	g at the date of hi	re for practice		3	Y	Ν	
	Documentation of education a	-		-					
4	 4 (dental health care personnel) and include hands on training for all staff assigned to process semi critical and critical instruments 3 Y 								
5								Ν	
6	Mechanism for corrective action	n for any deviatio	n from written po	licy. Documentation of any o	corrective actions	3	Y	Ν	
CON	CONFIDENTIAL EMPLOYEE HEALTH RECORDS								
7								Ν	
8	B Documentation of vaccinations offered to DHCP (Hepatitis B, Influenza), informed consent of exposure risk, and declinations of such vaccinations or immunizations 3							N	
9	Employee health records includes vaccination records						Y	Ν	
10	Employee health records kept for duration of employment, plus thirty years						Y	Ν	
11	Employee health records include any exposure and post exposure and follow up records						Y	Ν	
EXP	OSURE AND POST EXPOSUR								
12	2 Written policies and procedures regarding all occupational exposures which include a post exposure medical plan (e.g. use CDC needle stick/sharps injury/exposure protocol) 3						Y	Ν	
13								Ν	
14	Exposure and incident reporting forms						Y	Ν	
15	5 Sharps injury log							Ν	
MEDICAL CONDITIONS, WORK RELATED ILLNESS AND WORK RESTRICTIONS									
16							Y	Ν	
17	17Written policy and procedure for work restrictions for employees infected with or exposed to communicable diseases3						Y	N	
BLOODBORNE PATHOGEN ELEMENTS									
18	Written policies and procedures for the prevention of transmission of bloodborne pathogens3YN							Ν	
19	Written policies for hand hygiene, including documentation of training and appropriate selection of antiseptic agents 3 Y N							Ν	

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20	Written policies for use of personal protective equipment		3	Y	Ν				
21	Monitoring and documentation of compliance with PPE		3	Y	Ν				
22	Written policies and procedures for handling and management of sharps	3	Y	Ν					
	DISINFECTION AND STERLIZATION OF PATIENT CARE ITEMS								
23	Writen policies and procedures for managing semi-critical and critial items	3	Y	Ν					
	Written system outlining enitre sterilization process (written policies and procedures for transporting and	l							
24	processing of all contaminated critical and semi-critical instruments, the instrument processing area,		3	Y	Ν				
25	preparation and packaging of instruments, sterilization and storage of sterilized and clean dental instruments)								
25 26	Written policy and procedures for sterilization monitoring Weekly biological monitoring logs		3	Y Y	N N				
20	Current maintenance logs for sterilization equipment		3	Y	N				
			3	Y					
28	Weekly biological monitoring logs kept for 2+ years or since opening date:		3	Y	N N				
29	Written policy for managing failed chemical, heat or biological monitoring test		3	Y					
30	Equipment and manintenance logs IRONMENTAL INFECTION CONTROL ELEMENTS		3	T	N				
31	Written policy and procedure for aseptic management during patient care		3	Y	N				
31	Written policy and procedure for surface disinfection and environmental barrier protection			Y					
_			3	Y	N				
33 34	Written policy and procedure for medical waste management		3	Y	N				
_	Name/telephone number of licensed waste hauler for regulated waste				N				
35 36	Written Policy and procedure for decontaminating spills of blood or other body fluids				N N				
30	Written policy and procedure to improve dental unit water quality				N				
38	Documentation of dental unit water lines testing to meet potable water standard of EPA (<500 CFU/ml)				N				
30	Documentation of action taken to meet EPA potable water standard, including re-testing Written policy and procedure to maintain aesepis and prevent cross contamination when taking and								
39	processing dental radiographs				N				
40	Written policy and procedure to maintain aesepsis and prevent cross contamination during dental laboratory procedures				Ν				
ОТ⊦	IER								
41	A comprehensive and annually up-dated medical histroy form is used to evaluate patients								
COI	MMUNICABLE DISEASE CONTROL PROCEDURES	LEVEL 1-4	Y	N	N/A				
42	Single use or sterilization for critical items	1	Y	Ν	N/A				
43	Multi - dose vials used		Y	Ν					
44	a) if yes, vials are only entered with new, sterile syringe with a new, sterile needle 1				N/A				
45	b) Cap of multi-dose vial cleaned with alcohol based wipe before being accessed 2				N/A				
46	c) Are multi-use vials discarded when expired or 28 days after initial access (as applicable) - Must have date when first accessed	2	Y	Ν	N/A				
47	d) is initial access dated on the multi-use vials 2				N/A				
48	Fluid infusion and administration sets (IV bags, tubing and connectors) used?								
49	a) if yes, used only on one patient 1				N/A				
50	b) Disposed of after single use? 1			Ν	N/A				
51	c) Single IV bag is <u>not</u> used to mix medications for more than one patient 1			Ν	N/A				
52	d) Single dose medication/infusions are used for only one patient and discarded after use 1			Ν	N/A				
53	Personnel wear utility gloves when processing contaminated instruments - Not latex type for patient care 2								
54	Supplies for hand hygiene accessible to employees at point of need 2								
55	Soap and water easily accessible 2								
56	Alcohol based rubs easily accessible-if used	2	Y	N					
57	Team members display appropriate hand hygiene techniques	1	Y	Ν					

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A	PPROPRIATE PPE SUPPLIES ACCESSIBLE & EMPLOYEES WITH EXPOSURE RISKS				
58	Gloves (Latex and latex free or just latex free) Sterile Surgical Glovesfor surgical procedures	1/2	Y Y	N N	
59	Masks	1	Υ	Ν	
60	Safety glasses with side shield or full face shields	1	Y	Ν	
61	Disposable gowns/laundered gowns offered	1	Υ	Ν	
62	Health care workers display appropriate use of PPE barriers	2	Y	Ν	
63	Running water eye wash station accessible	3	Y	Ν	
64	Appropriate barrier products available (dental dams, protective eyewear, other)	2	Y	Ν	
65	Basic first aid products and equipment available (Recommended to include: nitrogylerin, Benadryl, epi-	4	Y	Z	
	pen, oxygen, aspirin, albuterol, glucose, glucagon)			IN	
DEN	TAL UNIT WATER QUALITY				
66	Dental unit water lines flushed between patients for a minimum of 20 seconds	2	Y	Ν	
67	Dental unit water lines are treated to remove biofilm	4	Y	Ν	
68	Maintain documentation of dental unit water line testing to meet the potable water standard of EPA (< 500 CFU/ml)	4	Y	Ν	
69	Maintain documentaion of dental unit water lines not meeting the potable water standard of EPA are treated and retested	4	Y	Ν	N/A
	CLEANING, DISINFECTION & STERILIZATION OF PATIENT CARE ITEMS				
70	Biofilm and organic matter are removed from critical and semi-critical instruments using detergents or enzymatic cleaners prior to sterilization	2	Y	Ν	
71	Sterilization equipment available and fully functional	1	Y	Ν	
72	Number of working autoclaves:	1	Y	Ν	N/A
73	Number of working chemiclaves:	1	Y	Ν	N/A
74	Number of working dry heat sterilizers:	1	Y	Ν	N/A
75	Number of working Flash steam sterilizers (Statim):	1	Y	Ν	N/A
76	Number of working ultrasonic cleaners:	1	Υ	Ν	
77	Endodontic files/instrumentation sterilized or disposed	1	Y	Ν	
78	Is Biological testing of sterilizer completed weekly	1	Y	Ν	
79	If independent biological testing service, Name:		Y	N	N/A
80	If in-office biological testing, is control processed?	2	Y	Ν	N/A
81	Sterilization cycles are verified with chemical/heat indicator. Both interior and external indicators	2	Y	Ν	
82	Critical items (any instrument that penetrates soft tissue or bone) instruments are sterilized after each use	1	Y	Ν	
	Use a biological indicator for every sterilizer load that contains a non-sterile Implantable device. Verify				
83	results before using the implantable device, whenever possible.	1	Y	Ν	N/A
84	Proper sterilization loading technique, not overloading	2	Y	Ν	
85	Heat Tolerant Handpieces are sterilized after each use (including high & low speed handpieces, prophylaxis angles, ultrasonic and sonic scaling tips, air abrasion devices, air and water syringe tips, and motorswith exception of electric type models)	1	Y	Ν	
86	Sterile packs are inspected for integrity, compromised packs are reprocessed	2	Y	Ν	
87	Event-related monitoring is used to monitor package integrity and packages are appropriately stored with a minimum of an initial date stamp	2	Y	Ν	
88	Single use instruments or devices are not processed and re-used	1	Y	Ν	
89	Semi-critical items are sterilized after each use if not heat sensitive	1	Y	Ν	
90	Heat sensitive semi-critical are at a minimum high level disinfected after each use or chemical sterilized after each use	1	Y	Ν	
91	Practice is using an FDA approved chemical sterilant	2	Y	N	N/A
92	All applicable label instruction are followed on EPA-registered chemical sterilant (dilution, shelf life, storage,	2	Y	N	N/A
93	safe use, disposal and material compatibility Practice is using an FDA approved high level disinfectant	2	Y	N	
93 94	Chemical used for high level disinfection are prepared according to manufacturer's instructions (dilution, shelf life, storage, safe use, disposal and material compatibility)	2	Y	N	
	ine, storage, sare ase, aisposar and material compatibility	1			

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95	Chemical used for high level disinfection are dated with expiration dates and discarded before expiration dates 2							
Asep	otic Techniques:							
96	Splash shields and equipment guards used on dental laboratory lathes	4	Y	Ν	N/A			
97	Fresh pumice and a sterilized, or new rag wheel used for each patient	2	Y	Ν	N/A			
98	Are devices used to polish, trim or adjust contaminated intraoral devices being disinfected or sterilized	2	Y	Ν	N/A			
99	Intraoral items such as impressions, bite registrations, prostheses and orthodontic appliances are cleaned and disinfected 2							
Envi	ronmental Infection Control		LEVEL 1-4	Y	Ν			
100	Semi-critical environmental surfaces (frequently touched surface that could potentially allow secondary transm to HCW or patients) are decontaminated between patients using a high level surface disinfectant	ission	2	Υ	Ν			
101	Noncritical environmental surfaces are decontaminated between patients		2	Y	Ν			
102	Objects and environmental surfaces are disinfected with an EPA registered tuberculocidal disinfectant at beginning of day							
103	Objects and environmental surfaces are disinfected with an EPA registered tuberculocidal disinfectant between patients							
104	Objects and environmental surfaces are disinfected with an EPA registered tuberculocidal disinfectant at the end of the day							
105	EPA registered tuberculocidal disinfectants are used at the dilution specified by the manufacturer							
106	All clinical contact surfaces are protected with barriers (especially areas that are difficult to clean)							
107	Clinical contact barriers are changed between patients							
108	Decontamination and clean areas separated in the instrument processing area							
109	Biohazardous waste is disposed of properly							
	Sharps							
110	Approved sharps containers utilized and accessible							
111	Sharps container taken out of service and processed appropriately							
112	Safe recapping techniques/devices used							
113	Sharps (needles, blades) are single use							
114	Employees use engineering controls (e.g., forceps) to retrieve contaminated sharps from trays or containers							

ACKNOWLEDGEMENT AND RECEIPT OF COPY BY OWNER/AUTHORIZED AGENT

The owner of the dental practice hereby acknowledges that by executing this document below and initialing each page's lower right hand corner on the line "Licensee Initials," receipt of a copy of this inspection/survey form is acknowledged.

In the event the dental practice has satisfactorily completed the inspection, as noted in this inspection/survey form, the owner/licensee will receive from the Board's Executive Director and/or representative, written notice of satisfactorily completing the inspection conducted.

If an owner/licensee has commenced the practice of dentistry prior to an Initial Inspection (NAC 631.1785) at any given location that inspection shall be deemed to be a Random Inspection pursuant to NAC 631.179.

If the inspection indicates "critical" deficiencies (items listed as "#1's") the owner/licensee will receive written notice from the Board's Executive Director and/or representative of the "critical" deficiencies and that a re-inspection will be conducted within seventy-two (72) hours of the written notice. However in the event the "critical" deficiencies noted, pose an immediate threat to the public health, safety and/or welfare the President of the Board, may without any further action of the Board, issue an Order of Summary Suspension pursuant to NAC 631.179(4).

In the event the inspection indicates "remedial action required" deficiencies (items listed as "#2's"), the owner/licensee will receive written notice from the Board's Executive Director and/or representative of the "remedial action required" deficiencies and that a re-inspection will be conducted within seven (7) days of the written notice.

In the event the inspection indicates "action required" deficiencies (items listed with a "#3"), the owner/licensee will receive written notice from the Board's Executive Director and/or representative of the "action required" deficiencies and that a re-inspection will be conducted within thirty (30) days of the written notice.

Receipt of a copy of the foregoing is hereby acknowledged;

Ву						Print name <u>:</u>
this	_day of	_, 20	at	<u> </u>	m.	Title and/or position/capacity: