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# Surgical preparation solution dry time: Summary

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#### **Specific Care Question**

For the patient who will be undergoing a surgical procedure, does the dry time of the products used for skin disinfection impact the risk of surgical site infection (SSI)?

#### **Recommendations Based on Current Literature**

No recommendation is can be made for or against optimal dry time of surgical site preparation agents, based on expert review of current literature by the Department of EBP. The overall certainty in the evidence is very low<sup>d</sup>. Only one randomized controlled study (Yasuda et al., 2015) was identified that compared surgical site preparation agents and there was no difference in site infections when comparing the group that did not have a wait time, to the group that had an approximate 5 minute wait time. When there is a lack of scientific evidence, standard work should be developed, implemented, and monitored (see Summary by Outcome for substantiation of this recommendation).

#### Literature Summary

**Background.** Skin antisepsis is the pre-operative treatment of intact skin in the operating room to reduce the microbial load on the patient's skin prior to making the surgical incision (WHO, 2018). Products used for skin antisepsis can be categorized as aqueous based or alcohol based (Armstrong, Patrick, & Erstad, 2001). Aqueous products are iodophor formulations, while alcohol-based products are formulations of isopropyl alcohol combined with iodophors or chlorhexidine (WHO, 2018). Alcohol based skin antiseptic agents are recommended for most surgeries (Berrios-Torres et al., 2017; WHO, 2018; AST, 2008), but these agents **should not** be used on mucous membranes (AST, 2008). Although alcohol-based products are valued for their quicker drying time (Armstrong et al., 2001; Magalini et al., 2013), they have been implicated in operating room fires (Jones et al., 2017; Weber, Hargunani, & Wax, 2006). This review will summarize identified literature on the topic.

**Study characteristics**. The search for suitable studies was completed on May 7, 2019. L. Harte, PharmD, CPHQ reviewed the seven titles and/or abstracts found in the search and identified six articles believed to answer the question. After an in-depth review one article, an RCT, answered the question (Yasuda et al., 2015). The Center for Disease Control (CDC) guidelines (Berrios-Torres et al. (2017) and the Association of Surgical Technologists (AST, 2008) are the primary source of information for this analysis. Additionally, one case study (Weber et al., 2006) and one *in vitro* study (Jones et al., 2017) were identified on risk of fire with solutions used for surgical preparation (prep).

#### Summary by Outcome

**Infection.** Yasuda et al. (2015) compared no wait time after application of povidone iodine (PVI) versus 5-minute wait time (approximate) after application of PVI and measured the outcomes of Positive Cultures and SSI (N = 89). The odds of having a positive culture were significantly less in the wait time group OR = 0.16, p = .008, 95% CI [0.04, 0.61]. For the outcome SSI, there was no difference in the number of SSI based on the wait time versus no wait time. There is very low certainty in this finding. The risk of bias is unclear, as randomization was not clearly reported, nor was blinding of subjects, personnel, or outcome assessors (see Figure 2). The evidence is indirect as only culture from the wound edge were reported, and a sample size calculation was not reported to know if enough subjects were recruited into the study. Finally, since only one study is included in this review, imprecision of the finding is serious (see Table 1).

**Other**. Although other trials were not identified that compared antiseptic dry time of various products with the outcome risk of SSI, the following points can be made:

- The CDC (Berrios-Torres et al., 2017) and the AST (AST, 2008) recommends that alcohol-based products be used for skin prophylaxis in preparation for surgery. However alcohol-based products should not be used on mucous membranes, rather aqueous iodophor products, such as PVI, are recommended for this surgery type (AST, 2008).
- Dry times (in seconds) of alcohol-based products and iodophor products are significantly less than dry times of iodophor products (see Figure 3):
  - ChlorPrep (alcohol based) vs. PVI (aquaeous based), *MD* = -53.0, 95% CI [-70.18, -35.82] (Magalini et al., 2013)



- DuraPrep (alcohol based) vs. PVI,(aquaeous based), MD = -31.8, 95% CI [-57.82, -5.78] (Armstrong et al., 2001)
- Weber et al. (2006) reported a case study of an operating room fire in a hirsute 62-year old male after surgical prep with DuraPrep. The operative field was draped after the patient's neck was shaved and the surgical prep solution was allowed to dry for at least 3 minutes. After skin incision and retraction, the electrocautery device was activated, and a flameless, smokeless fire occurred. Recommendations from this paper include:
  - Avoid the use of DuraPrep in the hirsute patient, collection of the agent on hair bearing skin can slow the dry time
  - The pooling of alcohol prep solution for any reason should be avoided
  - o Oxygen deliver during skin antisepsis should be a the minimal level to meet patient's need
- In an animal model, Jones et al. (2017) applied both alcohol (4% CHG with 70% isopropyl alcohol (IPA); plain IPA (70%); iodine-IPA, (0.7% iodine povacrylex and 74% IPA) and nonalcohol-based (4% CHG or 1% PVI paint) skin preps to porcine skin samples. Electrocautery was performed, with an electrosurgical pencil, immediately after application and after at least a 3-minute dry time.
  - Nonalcohol-based skin preps did not cause a fire for either dry times.
  - Alcohol-based skin preps did cause a fire in 22% (13/60) with no time allowed for prep to dry and 6% (10/60) where at least a 3-minute dry time was allowed.
  - Pooling of chlorhexidine-IPA created more fires
    - No time for prep dry there were 10% (2/20) fires in the no pooling group and 19/20 (95%) in the pooling group, p < .001
    - Time for prep to dry group there were 15% (3/20) fires in the no pooling group and 75 % (15/20) in the pooling group, p < .001

#### **Identification of Studies**

**Search Strategy and Results** (see Figure 1) PubMed: surgical site infection AND skin preparation AND (dry OR timing) Records identified through database searching n = 6Additional records identified through other sources n = 7

Studies Included in this Review

Citation	Study Type
Yasuda et al. (2015)	RCT

#### Studies Not Included in this Review with Exclusion Rationale

Citation	Reason for exclusion
Armstrong et al. (2001)	Does not address antiseptic drying time and SSI
AST (2008)	Does not address antiseptic drying time and SSI
Berrios-Torres et al. (2017)	Does not address antiseptic drying time and SSI
Hemani and Lepor (2009)	Does not address antiseptic drying time and SSI
Hibbard, Mulberry, and Brady (2002)	Does not address antiseptic drying time and SSI
Hibbard (2005)	Does not address antiseptic drying time and SSI
Johnson et al. (2016)	Does not address antiseptic drying time and SSI
Jones et al. (2017)	Does not address antiseptic drying time and SSI
Magalini et al. (2013)	Does not address antiseptic drying time and SSI
Moen, Noone, and Kirson (2002)	Does not address antiseptic drying time and SSI



	Solutions
Sidhwa and Itani (2015)	Does not address antiseptic drying time and SSI
Weber et al. (2006)	Does not address antiseptic drying time and SSI
WHO (2018)	Does not address antiseptic drying time and SSI
Methods Used for Appraisal and Synthesis	
	ation II (AGREE II) is an international instrument used to assess the quality and reporting of clinical practice
guidelines for this analysis (Brouwers et al.	2010).
<sup>b</sup> Rayyan is a web-based software used for the in 2017).	itial screening of titles and / or abstracts for this analysis (Ouzzani, Hammady, Fedorowicz & Elmagarmid,
<sup>c</sup> Review Manager (Higgins & Green, 2011) is a C and create the forest plots found in this and	Cochrane Collaborative computer program used to assess the study characteristics as well as the risk of bias lysis.
	$\dot{\mathbf{D}}$ is the tool used to create the Summary of Findings table(s) for this analysis (see Table 1).
<sup>e</sup> The Preferred Reporting Items for Systematic R screened, and eligibility criteria is applied (M	eviews and Meta-Analyses (PRISMA) flow diagram depicts the process in which literature is searched, Joher, Liberati, Tetzlaff, & Altman, 2009).
	S Consortium. (2010) AGREE II: Advancing guideline development, reporting and evaluation in Journal, 182, E839-842. Retrieved from <u>https://www.agreetrust.org/wp-content/uploads/2017/12/AGREE-</u> t-2009-Update-2017 pdf
	Elmagarmid, A. (2016). Rayyan-a web and mobile app for systematic reviews. Systematic Reviews, 5(1),
	ne Handbook for Systematic Reviews of Interventions [updated March 2011] (Version 5.1.0 ed.): The
<sup>d</sup> GRADEpro GDT: GRADEpro Guideline Developm from <u>gradepro.org</u> .	nent Tool (2015). McMaster University, (developed by Evidence Prime, Inc.). [Software]. Available
<sup>e</sup> Moher D, Liberati A, Tetzlaff J, Altman DG, The	PRISMA Group (2009). <i>P</i> referred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA oi:10.1371/journal.pmed1000097 For more information, visit <u>www.prisma-statement.org</u> .
Question Originator	
Lory Harte, PharmD, CPHQ	
Medical Librarian Responsible for the Searc	h Strategy
Keri Swaggart, MLIS, AHIP	
EBP Scholar's Responsible for Analyzing the Erin Lindhorst, MS, RD, LD	e Literature
	ing, Synthesizing, and Developing this Document
Nancy Allen, MS, MLS, RD, LD, CPHQ	ng, synthesizing, and beveloping this becament
Acronyms Used in this Document	
Acronym Explanation	
AST Association of Surgical Techno	plogists
CAT Critically Appraised Topic	
CDC Centers for Disease Control	
CHG Chlorhexidine	
CMH Children's Mercy Hospital	
CDC Clinical Duration Cuidaling	

CPG Clinical Practice Guideline



EBP	Evidence Based Practice
IPA	Isopropyl Alcohol
MD	Mean Difference
OR	Odds Ratio
PRISMA	Preferred Reporting Items for Systematic Reviews and Meta-Analysis
Prep	Preparation
PVI	Povidone Iodine
RCT	Randomized Controlled Trial
SSI	Surgical Site Infection
WHO	World Health Organization
Date Develo	ped/Updated
July 2019	



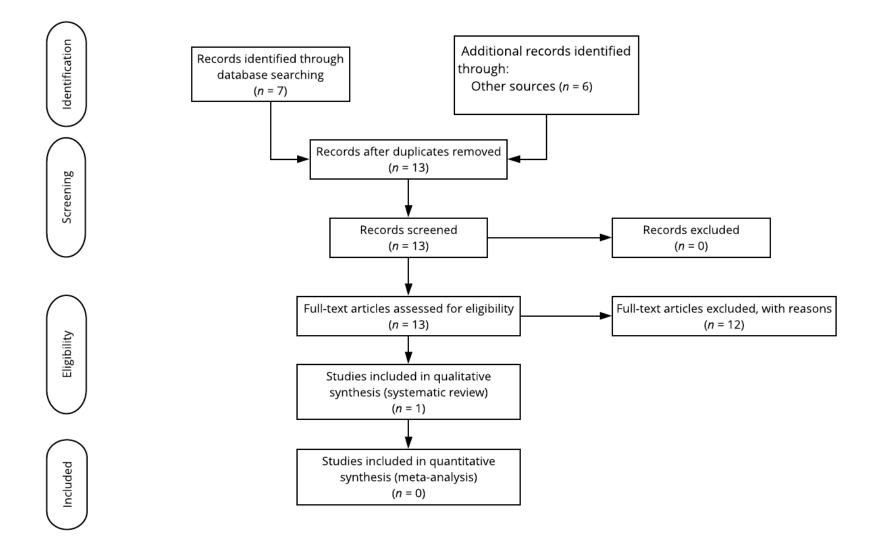


Figure 1. Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRIMSA)<sup>e</sup>

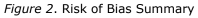


AGREE II<sup>a</sup> Summary for the CDC Prevention of SSI Guideline (Berrios-Torres et al., 2017)

Domain	Percent Agreement			
Scope and purpose	86%			
Stakeholder involvement	81%			
Rigor of development	81%			
Clarity and presentation	93%			
Applicability	46%			
Editorial independence	94%			
Overall guideline assessment	90%			
Team's recommendation for guideline use Yes with modification				
Note: Four EBP Scholars completed the AGREE II on this guideline.				









#### Table 1

#### Summary of Findings Table: Wait Time vs. No Wait Time for Surgical Site Preparation Solution

	Certainty assessment				Summary of findings						
Nº of						Overall	Study event rates (%)			Anticipated absolute effects	
participants (studies) Follow-up	rticipants of Inconsistency Indirectness Imprecision Publication certainty bias of	With no wait time	With Wait time	<ul> <li>Relative effect (95% CI)</li> </ul>	Risk with no wait time	Risk difference with Wait time					
Positive Cu	ulture	S									
89 (1 RCT)	serious <sup>a</sup>	not serious	serious <sup>b</sup>	serious <sup>c</sup>	none	⊕⊖⊖⊖ VERY LOW	13/43 (30.2%)	3/46 (6.5%)	<b>OR 0.16</b> (0.04 to 0.61)	302 per 1,000	<b>237 fewer</b> <b>per 1,000</b> (from 285 fewer to 93 fewer)
SSI Infect	ion										
89 (1 study)	serious ª	not serious	serious <sup>b</sup>	serious <sup>c</sup>	none	-	0/43 (0.0%)	2/46 (4.3%)	<b>OR 4.89</b> (0.23 to 104.76)	0 per 1,000	0 fewer per 1,000 (from 0 fewer to 0 fewer)

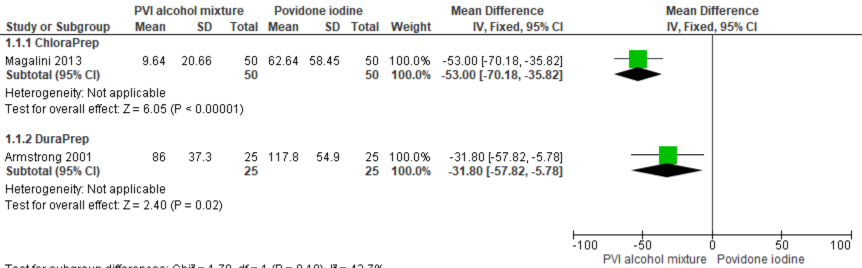
Notes:

a. Reporting on randomization; allocation concealment; blinding of participant, personnel, and outcome assessors is poorly reported;

b. Cultures were only collected from the wound edge;

c. It is a single study with 89 subjects.





Test for subgroup differences: Chi<sup>2</sup> = 1.78, df = 1 (P = 0.18), l<sup>2</sup> = 43.7%

Figure 3. Comparison of PVI alcohol mixture vs. PVI/iodophor, Outcome: Dry time



Characteristics of Studies

#### Armstrong 2001

Methods	Clinical Trail (not randomized)
Participants	<ul> <li>Participants: <ul> <li>Twenty-five operating room personnel</li> <li>Twenty-five subjects (patient volunteers)</li> </ul> </li> <li>Setting: College of Pharmacy, Arizona, US</li> <li>Participated in study: N = 50 <ul> <li>Group 1, Operating room personnel*: n = 25</li> <li>Povidone iodine paint and scrub (7.5% povidone iodine, 10% water, Operand; APlicare, Inc., Branford CT</li> <li>Duraprep (0.7% iodophor, 74% isopropyl alcohol; 3M Health Care plus Ioban in combination</li> <li>Prevail (5% povidone iodine, 62% alcohol; Allegiance Health care Corp., McGaw Park, IL</li> <li>LiquiDrape not FDA approved at time of the study, Tradmark for this product has been abandoned https://trademark.trademarkia.com/liquidrape-75404162.html July 2 2019</li> <li>Group 2, Patient volunteers: n = 25</li> <li>Group 1, Applied surgical prep *: n = 25</li> <li>Group 1, Applied surgical prep *: n = 25</li> <li>Group 2, Had surgical prep applied: n = 25</li> <li>Gender, males (as defined by researchers): <ul> <li>Not reported</li> </ul> </li> <li>Race / ethnicity or nationality (as defined by researchers): <ul> <li>Not reported</li> </ul> </li> <li>Both groups - greater than or equal to 18 years of age</li> <li>Both groups - greater than or equal to 18 years of age</li> <li>Both groups - free of known hypersensitivity to povidone iodine or alcohol</li> <li>Operating room personnel - at least six months experience assisting in pre-operative patient preparation</li> </ul> </li> </ul>
Interventions	<ul> <li>Operating room personnel applied the four skin prep formulations to the lower extremity of the patient volunteers. There were two rounds of application. The first round, two products were applied, one to each leg from the knee to the ankle. The product was allowed to dry and then removed. The process was repeated with the remaining skin prep products for the second round.</li> <li>Patients were not required to shave their legs</li> <li>Operating room personnel read the product insert instructions prior to application.</li> </ul>



Outcomes	Primary outcome(s):			
	Product application			
	Drying time			
	Removal time			
	Overall satisfaction			

#### Magalini 2013

Methods	Observational study
Participants	Participants: Surgeons performing elective and emergency surgeries (medium and major operations).         Setting: Hospital, Italy         Number enrolled into study: N = 100         • Group 1, Povidone iodine (PVI): n = 50         • Group 2, ChloraPrep: n = 50         Number completed: N = 100         • Group 1: n = 50         • Group 2: n = 50         Gender, males: Not reported         Race / ethnicity or nationality (as defined by researchers):         • Not reported         Age: Not reported         Inclusion criteria:         • Surgeon approval for observation         • Use of either PVI or ChloraPrep         Exclusion criteria:         • None listed         Covariates identified: not reported
Interventions	<ul> <li>Both: Every surgeon (27 unique surgeons were observed) performed their own surgical field and uses the two different approaches defined below they start from the middle of the surgical field and swabbing out. When the field is almost dry they may use a paper towel to complete the drying. All surgeons identified that they had received product training.</li> <li>Group 1: PVI is poured on the skin and gauze/clamp used</li> <li>Group 2: ChloraPrep applicator</li> </ul>
Outcomes	<ul> <li>Primary outcome(s):         <ul> <li>Comparison of IPV and ChloraPrep in supplies used</li> <li>Comparison of IPV and ChloraPrep in time for application, drying*, and total time needed for disinfection (defined as from the beginning of painting to placing of drapes)</li> </ul> </li> <li>Secondary outcome(s)         <ul> <li>Surgeons opinion from the questionnaire</li> <li>Youtcomes of interest to the CMH CPG or CAT development team</li> </ul> </li> </ul>



#### Yasuda 2014

Methods	Prospective, Randomized, controlled study
Participants	Setting: Department of Orthopaedic Surgery, Hamamatsu University of Medicine         Randomized into study: N = 89         • Group 1: No wait time for povidone-iodine applied, n = 43         • Group 2: Wait time povidone-iodine applied, n = 46         Completed study: N = 89         • Group 1: n = 43         • Group 1: n = 43         • Group 1: n = 43         • Group 2: n = 46         Gender, males:         • Group 1: n = 21         • Group 2: n = 23         Race / ethnicity or nationality (as defined by researchers):         • Not reported         Age, years (mean):         • Group 1: 61.9         • Group 2: 58.1         Inclusion criteria:         • Patients scheduled for spinal surgery         Exclusion criteria:         • Not reported         Power Analysis:         • Not reported
Interventions	<ul> <li>Both:         <ul> <li>In all cases, the surgical field was sealed with an antimicrobial plastic adhesive wound drape just before starting the surgery.</li> <li>Culture samples were collected by rubbing a cotton swab at the wound edge just before wound closure and then they were incubated at 37-degree Celsius for 5 to 7 days.</li> <li>Cefazolin was administered three times on the day of surgery, before surgery, one hour after surgery, and six hours after surgery, and two times on the next day as a prophylactic antibiotic.</li> </ul> </li> <li>Group 1: povidone-iodine was applied to the surgical site just before skin incision, after the surgeon's hands were scrubbed.</li> <li>Group 2: povidone-iodine was applied before the surgeon's hands were scrubbed. Expected Wait time 5 minutes.</li> </ul>
Outcomes	Primary outcome: • Culture results Secondary outcome: • SSI infection
Results	• In Group 1, coagulase negative <i>Staphylococcus aureus</i> was identified in one culture. In Group 2, three different bacteria ( <i>streptococcus, staphylococcus epidermidis</i> , and <i>coagulase negative staphylococcus</i> ) were identified in the culture.



<ul> <li>Two cases of SSI (deep infection) (2 out of 46 patients, 4.3%) were identified in group 2 four weeks after surgery, and cultures from the wound edge intraoperatively were negative. There was no case of SSI in Group 1 after the surgery.</li> </ul>
• Because bacteria on the skin appeared significantly reduced by allowing povidone-iodine to dry for several minutes prior to surgery, the researchers recommend this approach to reduce the incidence of postoperative infections. The recommended drying time prior to surgery is 10 minutes.
<ul> <li>A limitation of this study is that only analysis of cultures from the wound edge was conducted.</li> </ul>

Risk of bias table

Bias	Scholars' judgment	Support for judgment	
Random sequence generation (selection bias)	Unclear risk	Patients were randomly allocated into 2 groups, however how they were randomized was not indicated.	
Allocation concealment (selection bias)	Unclear risk	Insufficient information to determine	
Blinding of participants and personnel (performance bias)	Unclear risk	The study did not address this outcome	
Blinding of outcome assessment (detection bias)	Unclear risk	Insufficient information to determine	
Incomplete outcome data (attrition bias)	Unclear risk	All patients enrolled were analyzed. Although, no power analysis performed	
Selective reporting (reporting bias)	Unclear risk	SSI infections not tested for significance	
Other bias	Low risk	The study appears to be free of other sources of bias.	



#### References

- Armstrong, E. P., Patrick, K. L., & Erstad, B. L. (2001). Comparison of preoperative skin preparation products. *Pharmacotherapy*, *21*(3), 345-350. Retrieved from https://www.ncbi.nlm.nih.gov/pubmed/11253858
- Berrios-Torres, S. I., Umscheid, C. A., Bratzler, D. W., Leas, B., Stone, E. C., Kelz, R. R., . . . Healthcare Infection Control Practices Advisory, C. (2017). Centers for Disease Control and Prevention Guideline for the Prevention of Surgical Site Infection, 2017. *JAMA Surg*, 152(8), 784-791. doi:10.1001/jamasurg.2017.0904
- Hemani, M. L., & Lepor, H. (2009). Skin preparation for the prevention of surgical site infection: which agent is best? *Rev Urol, 11*(4), 190-195. Retrieved from https://www.ncbi.nlm.nih.gov/pubmed/20111631
- Hibbard, J. S. (2005). Analysis comparing the antimicrobial activity and safety of current antiseptic agents: A review. Journal of Infusion Nursing, 16(2), 195.
- Hibbard, J. S., Mulberry, G. K., & Brady, A. R. (2002). A clinical study comparing the skin antisepsis and safety of ChloraPrep, 70% isopropyl alcohol, and 2% aqueous chlorhexidine. *J Infus Nurs*, 25(4), 244-249. Retrieved from https://www.ncbi.nlm.nih.gov/pubmed/12131506
- Johnson, M. P., Kim, S. J., Langstraat, C. L., Jain, S., Habermann, E. B., Wentink, J. E., . . . Bakkum-Gamez, J. N. (2016). Using Bundled Interventions to Reduce Surgical Site Infection After Major Gynecologic Cancer Surgery. *Obstet Gynecol*, *127*(6), 1135-1144. doi:10.1097/AOG.00000000001449
- Jones, E. L., Overbey, D. M., Chapman, B. C., Jones, T. S., Hilton, S. A., Moore, J. T., & Robinson, T. N. (2017). Operating Room Fires and Surgical Skin Preparation. J Am Coll Surg, 225(1), 160-165. doi:10.1016/j.jamcollsurg.2017.01.058
- Magalini, S., Pepe, G., Panunzi, S., De Gaetano, A., Abatini, C., Di Giorgio, A., . . . Gui, D. (2013). Observational study on preoperative surgical field disinfection: povidone-iodine and chlorhexidine-alcohol. *Eur Rev Med Pharmacol Sci*, *17*(24), 3367-3375.
- Moen, M. D., Noone, M. B., & Kirson, I. (2002). Povidone-iodine spray technique versus traditional scrub-paint technique for preoperative abdominal wall preparation. *Am J Obstet Gynecol, 187*(6), 1434-1436; discussion 1436-1437.
- Organization, W. H. (2018). *Global guidelines on the prevention of surgical site infection*. WHO Retrieved from https://www.who.int/gpsc/appendix8.pdf Sidhwa, F., & Itani, K. M. (2015). Skin preparation before surgery: options and evidence. *Surg Infect (Larchmt), 16*(1), 14-23. doi:10.1089/sur.2015.010 Association of Surgical Technologists. (2008). AST standards of practice for skin prep of the surgical patient. Retrieved from
  - http://www.ast.org/uploadedFiles/Main\_Site/Content/About\_Us/Standard\_Skin\_Prep.pdf
- Weber, S. M., Hargunani, C. A., & Wax, M. K. (2006). DuraPrep and the risk of fire during tracheostomy. *Head Neck, 28*(7), 649-652. doi:10.1002/hed.20396 Yasuda, T., Hasegawa, T., Yamato, Y., Kobayashi, S., Togawa, D., Arima, H., & Matsuyama, Y. (2015). Optimal Timing of Preoperative Skin Preparation with Povidone-Iodine for Spine Surgery: A Prospective, Randomized Controlled Study. *Asian Spine J, 9*(3), 423-426. doi:10.4184/asj.2015.9.3.423

