Journal of Epidemiology and Global Health (2012) 2, 173-17





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Hand rub dose needed for a single disinfection varies according to product: A bias in benchmarking using indirect hand hygiene indicator

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Received 10 August 2012; received in revised form 9 October 2012; accepted 10 October 2012 Available online 20 November 2012

KEYWORDS

Hand hygiene; Compliance; Hand rub; Benchmarking **Abstract** *Background:* The 3 ml volume currently used as the hand hygiene (HH) measure has been explored as the pertinent dose for an indirect indicator of HH compliance. A multicenter study was conducted in order to ascertain the required dose using different products.

Method: The average contact duration before drying was measured and compared with references. Effective hand coverage had to include the whole hand and the wrist. Two durations were chosen as points of reference: 30 s, as given by guidelines, and the duration validated by the European standard EN 1500. Each product was to be tested, using standardized procedures, by three nosocomial infection prevention teams, for three different doses (3, 2 and 1.5 ml).

Results: Data from 27 products and 1706 tests were analyzed. Depending on the product, the dose needed to ensure a 30-s contact duration in 75% of tests ranging from 2 ml to more than 3 ml, and to ensure a contact duration exceeding the EN 1500 times in 75% of tests ranging from 1.5 ml to more than 3 ml. The aftermath interpretation is the following: if different products are used, the volume utilized does not give an unbiased estimation of the HH compliance. Other compliance evaluation methods remain necessary for efficient benchmarking.

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1. Introduction

With the development of benchmarking and public reporting for infection control, indirect hand hygiene (HH) indicators have been developed, with the aim of giving repeated data more frequently than was possible with classic compliance studies [1]. The most common HH indicator translates a quantity of consumed hand rubs into a number of HH actions and calculates the number of HH actions by patients/days. This method has been successfully used for time series analysis on the evaluation of prevention policies or for comparison between units within a hospital or within a hospital group using the same products [2-6], but some studies have highlighted the limited use of this tool for wider comparisons [7–9] with an exception in the case of a very large difference [7]. The necessity to validate each component of such an indicator has arisen (i.e., dose, activity reference) for effective reproducibility and unbiased historic comparisons. The European historic choice of a 3 ml dose as reference is based on the EN 1500 standard, which required the comparison of tested products to two applications of 3 ml of propanol [10]. At the present time, this dose is challenged because hand rub suppliers do recommend different doses on their instructions and have also validated the standard EN 1500 with different doses. Furthermore, there is no systematic link between the suppliers' recommended dose and the one needed to validate the EN 1500. The actual dose distributed by cheap dispensers is frequently variable as well. The initial dose ranges from 1 to 3 ml, but after a short period of use, the actual dose is very frequently smaller as demonstrated by the Kohan study [11]. The aim of this multicenter prospective open trial was to define the required dose for correct HH while using different rubs.

2. Methods

2.1. Objective

For a dose and a product, the evaluation criteria were: proper coverage of the whole hand, average contact duration before drying, and accuracy of this duration according to available references. Three doses were tested (3, 2 and 1.5 ml).

2.2. Definition of correct duration of hand hygiene

For the study, the duration of HH was considered correct for a dose and a product if:

- According to the 2002 French recommendations (http://www.sf2h.net/publications-SF2H/SF2H_ recommandations_hygiene-des-mains-2009.pdf), at least 75% of tests allowed for contact duration longer than or equal to 30 s.
- According to the EN 1500, at least 75% of tests allowed for contact duration longer than or equal to that of the contact time validated by the product by means of EN 1500 [10].

Duration was counted in seconds from the spreading of the product to the very moment when any treated part of the hand is considered dry. A horizontal light was used for better evaluation. The dose was measured with a syringe (one syringe was used for every product). Duration was measured by the operator with a stopwatch.

2.3. Definition of common hand hygiene procedure

The selected procedure included the six steps which were described in EN 1500, completed by a wrist rubbing. While this step is not included in EN 1500, as there are no microbiological references to prove its effectiveness, it is still recommended in French health care facilities, and it should be taken into account for consumption measurement. This selected procedure was used in all test centers. A training course was organized in order to harmonize the procedures of this study.

2.4. Selection of products

As products had to meet the French requirements, 59 products were selected and their suppliers contacted in order to supply free samples. They were chosen from the 2008 "Liste Positive Désinfectants (LPD)," a publication of the French Society of Hospital Hygiene (http://www.sf2h.net/publications-SF2H/SF2H_LPD-2009.pdf), which presented the disinfectants conforming to a list of efficacy tests, and from the "Liste Positive Désinfectants Dentaires," which was the equivalent publication for use by dentists (http://sf2h.net/publications-SF2H/SF2H-ADF_LPDdentaire-2009.pdf). products that were marketed too recently to be accepted by the LPD group were also included in the selection. All free available products were tested. The order between products was randomly assigned in each center.

2.5. Test centers

Five test centers were set up: Bordeaux, Lille, Lyon, Rennes and Strasbourg. Each person taking part in the test was an infection control worker and each product had to be tested for each dosage

(3, 2 and 1.5 ml) in three centers and at least seven times per center and per dosage.

2.6. Organization and analysis

The tests were carried out between May and July 2008. For each dose, if a product had not been tested 10 times, it was then taken out of the analysis. The data were analyzed with Epi Info 2002 and SPSS V12. The duration of hand rubbing was considered correct according to the 2002 French guidelines, if this value is longer than or equal to 30 s, and effective according to the EN 1500 and if this value is longer than or equal to that of the contact time validated by the product by means of EN 1500.

3. Results

During the study, 1800 tests were performed. Between 9 and 111 tests were carried out per product. The number of professionals participating in the testing process was 71 (21 men and 50 women). Only 27 products and 1706 tests were included in the analysis, as four scheduled products were not available in three test centers and were tested less than 10 times. The included products were Actisène alcogel (Werner and Mertz), Alco Aloe solution and gel (Ansell), Alcocide, Alcoogel H and Septigel (Prodene Klint), Anios gel 85 NPC and Manugel 85 NPC (Anios), Assanis pro (Blue Skin SA), Dermalkan (Alkapharm), Desderman gel and Desderman N (Schülke), Elusept gel and Elusept solution (Elusept), GHA (Cellande), Manupure (Elis), Manurub and Manurub gel (Steridine), Nosocomia gel (Prodene Klint), Procide (IPC), Purell and Purell 85 (Gojo), Softalind hand sanitizer (BBraun), Spirigel and Spitacid (Ecolab), Sterilium (Rivadis), and Stoko progel and Stokosept (Stoko).

3.1. Results per dose and per product

For a 1.5 ml dose, the dose was sufficient to cover both hands and wrists in 551/575 cases (95.8%). Variability between the average contact duration before drying of each product was very important ($p < 10^{-3}$) (Fig. 1A and Table 1). According to the 30-s duration, there was not one single product for which 75% of tests showed a drying duration longer than or equal to the reference, but, according to EN 1500 there were 3/27 tested products for which 75% of tests showed a drying duration longer than or equal to the validated standard duration: Purell, Purell 85 and Softalind hand sanitizer, which are validated for the 15-s duration.

For a 2 ml dose, the dose was sufficient to cover both hands and wrists in 530/538 cases (98.5%). Variability between the average contact duration before drying of each product was very important $(p < 10^{-3})$ (Fig. 1B and Table 1). According to the 30-s duration, there were 5/26 tested products for which 75% of tests showed a drying duration longer than or equal to the reference: GHA, Manurub, Purell, Sterillium and Stokosept gel. According to EN 1500, there were six products for which 75% of tests showed a drying duration longer than or equal to the validated standard duration: all products which were validated for the 15-s duration (Purell, Purell 85, Softalind hand sanitizer), and 3 products which were validated for the 30-s duration (GHA, Manurub and Sterillium).

For a 3 ml dose, the dose was sufficient to cover both hands and wrists in 592/593 cases (99.8%). Variability between the average contact duration before drying of each product was very important ($p < 10^{-3}$) (Fig. 1C and Table 1). According to the 30-s duration, all 26 tested products, except Softalind hand sanitizer, showed a drying duration longer than the reference. According to EN 1500, for 18 products, 75% of tests showed a drying duration longer than the expected duration defined by EN 1500. This criterion was not validated for all products that passed the standard 30 s twice, repeatedly.

4. Discussion

In this study, correct coverage of hands and wrists was achieved for all tested products and all doses. Depending on the product, the dose needed to ensure a contact of 30 s, in 75% of the tests, varied from 2 ml to more than 3 ml. The dose needed to ensure the requisite contact duration for the validation of the standard EN 1500 in 75% of the tests ranged from 1.5 ml to more than 3 ml. For some products, the dose needed was higher than 3 ml (the maximum dose tested), consequently it was not checked.

This result is consistent with previous observations during local tests which showed that, within the same unit and for the same test duration, required doses varied between products [12]. Consequently, an estimation of the compliance (number of HH made divided by the number of necessary HH) by the quantity of product used (in ml per patient day, for example) is not accurate, and a comparison between hospitals could be biased, independent regardless of the limits of the prediction of the necessary HH by the type of service and per patient days. It is pertinent within a hospital

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where units use the same product and over a period of time if this product remained unchanged, but seems biased for a national score [4].

This study considers it important to assess the representativeness and reproducibility of the study. A large number of marketed products were tested in five different centers, but without randomization or exhaustiveness, it is impossible to

extend results to all marketed products or to all hospitals for this matter. The tests were carried out over the same period, and it does not seem relevant to take into account any seasonal effect, which is a well-known confounding factor in tolerance studies and professional conduct. The variation between required doses according to the product may be associated with the consistency

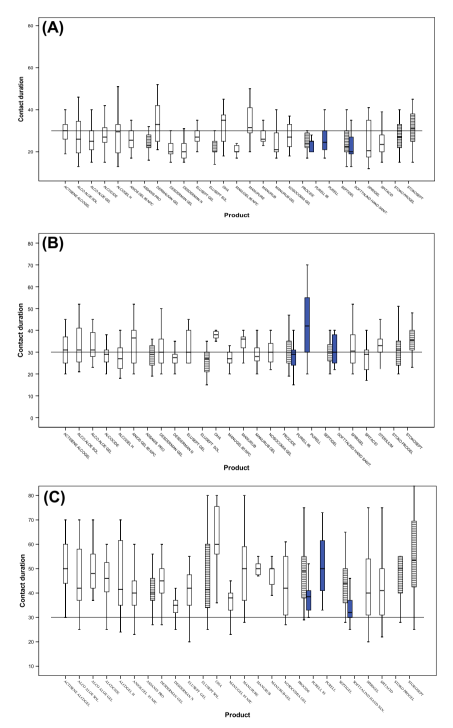


Figure 1 Contact hand rubbing duration in seconds; distribution by product and by dose. EN 1500 validation: ■ 15 s; □ 30 s; ■ 60 s. (A) For a 1.5 ml dose; (B) for a 2 ml dose; and (C) for a 3 ml dose.

Table 1 Drying durations by dose and product: value of percentile 25 of duration of tests made for each product and each duration (i.e. 75% of tests have been longer than this value).

EN 1500 contact time validated ^a	Products	Dose 1.5 ml		Dose 2 ml		Dose 3 ml	
		Tests number	Percentile 25 ^b (s)	Tests number	Percentile 25 ^b (s)	Tests number	Percentile 25 ^b (s)
15 s	Purell	28	21	33	30	28	41
	Purell 85	14	20	21	24	14	32
	Softalind hand sanitizer	14	19	14	25	14	28
30 s	Actisène alcoogel	25	26	14	25	30	44
	Alcocide	19	24	19	25	19	40
	Alcogel H	24	19	24	22	24	35
	Anios gel 85 NPC	22	22	26	25	29	35
	Dermalkan	14	25	_	_	_	_
	Desderman gel	41	19	28	25	42	40
	Desderman N	32	17	30	25	34	32
	Elusept gel	11	25	13	25	11	30
	GHA	12	25	11	35	12	56
	Manugel 85 NPC	21	20	31	25	21	33
	Manupure	14	28	_	_	14	37
	Manurub	18	25	17	31	19	47
	Manurub gel	19	20	7	28	27	43
	Nosocomia gel	19	22	19	25	19	31
	Spitacid	14	20	14	22	14	30
	Spirigel	28	17	28	25	28	31
	Sterilium	_	_	15	30	_	_
	Alco Aloe gel	26	21	33	28	26	42
	Alco Aloe solution	27	19	14	19	25	21
$2 \times 30 \text{ s}$	Assanis pro	21	22	21	24	21	37
	Elusept solution	25	18	19	20	26	30
	Procide	14	22	14	25	15	38
	Septigel	36	20	24	26	34	36
	Stoko progel	17	22	20	24	14	39
	Stokosept	20	25	24	31	32	41

^a Suppliers data.

(liquid or gel, alcohol concentration, various additives), but no systematic association was to be found in this study, and this point is debated [13,14]. This study does not give correcting factors, and other measures are necessary with this in mind. The bias of the large difference, in the field, on habits of rubbing techniques, on the actual dose and of hand size have been eliminated by the standardization of the method, the use of a measured dose and the repartition of the test between the different centers.

If one wanted to define the ideal dose needed for accurate proper hand rub disinfection, the protocol should take into account all confounding factors. The choice of a pertinent reference is also necessary. This study used the EN 1500 standard, as required in Europe for all marketed products, as proof of a real efficacy on bacteria in the case

of hygienic hand disinfection, but the duration of 30 s was equally used, as the required duration for a minimal activity on viruses. When the ISO system includes similar standards, it will be more adapted for international comparison, some products being available in many countries. It is important, too, to perform a study on a sample of sufficient size, in a representative selection of healthcare facilities and climates. But such a test could give more data on products, with the aim of eliminating the effect of the product in comparisons based on consumption measures.

The range of available hydro-alcoholic solutions, gels or foams has considerably increased over the last few years. Hand rub is now the main HH method, and indirect monitoring of HH compliance, easier than observational studies, is frequently used. But it is only useful if complete data are available on

^b Value of percentile 25: the duration of hand rubbing was considered effective, according to the 2002 French guidelines, if this value is longer than or equal to 30 s, and effective according to the EN 1500 if this value is longer than or equal to that of the contact time validated by the product by means of EN 1500.

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consumption and activity by unit, aggregated data at an entire hospital being of little use in the field.

Indeed, in the field of HH evaluation, it is necessary to use indicators for benchmarking or for evaluation of campaigns [15]. All indicators could be biased: HH compliance measured by observation could also be biased if different methods of observation or analysis are used [16]. Local customs can also interfere: interpretation of five moments of the WHO guidelines seems to differ between services, for example, or encouraging the use of products by patients and visitors can increase consumption [17]. The only safeguards against all these biases are the continuous use of the same method or the introduction of a correction in comparison, and the association of different media: HH observation, hand rub consumption or new techniques, such as electronic devices [18,19].

In conclusion, the old idea of an exact relationship between the volume of hand rub consumed and HH compliance remains false, and necessary comparisons should be made with care. Further studies appear necessary, in order to develop tools for modulating this relationship, while remaining circumspect about the limits of the use of the volume of product used as an indicator of good compliance.

Acknowledgements

To Marie Yvonne Dixon and Ian Russell for English text revision, and to product suppliers for products. No grant was obtained for this study. No conflict of interest to declare.

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