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Enhanced terminal room disinfection and acquisition and infection caused by multidrug-resistant organisms and *Clostridium difficile* (the Benefits of Enhanced Terminal Room Disinfection study): a cluster-randomised, multicentre, crossover study

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Summary

Background—Patients admitted to hospital can acquire multidrug-resistant organisms and *Clostridium difficile* from inadequately disinfected environmental surfaces. We determined the effect of three enhanced strategies for terminal room disinfection (disinfection of a room between occupying patients) on acquisition and infection due to meticillin-resistant *Staphylococcus aureus*, vancomycin-resistant enterococci, *C difficile*, and multidrug-resistant *Acinetobacter*.

Methods—We did a pragmatic, cluster-randomised, crossover trial at nine hospitals in the southeastern USA. Rooms from which a patient with infection or colonisation with a target organism was discharged were terminally disinfected with one of four strategies: reference

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See **Online** for appendix

Contributors

DJA did the literature search, prepared the figures, designed the study, collected, analysed, and interpreted data, wrote the Article, and approved the final version. LFC, DJW, RWM, SSL, WAR, and DJS designed the study, collected and interpreted data, revised the Article, and approved the final version. PFT, MB, PB, JCS, LPK, HK, and MGT collected data, revised the Article, and approved the final version. YL analysed and interpreted data, revised the Article, and approved the final version.

Declaration of interests

WAR and DJW have received consulting fees from Clorox. The other authors declare no competing interests.

(quaternaly ammonium disinfectant except for *C difficile*, for which bleach was used); UV (quaternaly ammonium disinfectant and disinfecting ultraviolet [UV-C] light except for *C difficile*, for which bleach and UV-C were used); bleach; and bleach and UV-C. The next patient admitted to the targeted room was considered exposed. Every strategy was used at each hospital in four consecutive 7-month periods. We randomly assigned the sequence of strategies for each hospital (1:1:1:1). The primary outcomes were the incidence of infection or colonisation with all target organisms among exposed patients and the incidence of *C difficile* infection among exposed patients in the intention-to-treat population. This trial is registered with ClinicalTrials.gov, NCT01579370.

Findings—31 226 patients were exposed; 21 395 (69%) met all inclusion criteria, including 4916 in the reference group, 5178 in the UV group, 5438 in the bleach group, and 5863 in the bleach and UV group. 115 patients had the primary outcome during 22 426 exposure days in the reference group (51.3 per 10 000 exposure days). The incidence of target organisms among exposed patients was significantly lower after adding UV to standard cleaning strategies (n=76; 33.9 cases per 10 000 exposure days; relative risk [RR] 0.70, 95% CI 0.50–0.98; p=0.036). The primary outcome was not statistically lower with bleach (n=101; 41.6 cases per 10 000 exposure days; RR 0.85, 95% CI 0.69–1.04; p=0.116), or bleach and UV (n=131; 45.6 cases per 10 000 exposure days; RR 0.91, 95% CI 0.76–1.09; p=0.303) among exposed patients. Similarly, the incidence of *C difficile* infection among exposed patients was not changed after adding UV to cleaning with bleach (n=38 vs 36; 30.4 cases vs 31.6 cases per 10 000 exposure days; RR 1.0, 95% CI 0.57–1.75; p=0.997).

Interpretation—A contaminated health-care environment is an important source for acquisition of pathogens; enhanced terminal room disinfection decreases this risk.

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Introduction

Multidrug-resistant organisms and *Clostridium difficile* are common causes of health-care-associated infections that lead to adverse patient outcomes.¹ The hospital environment may be an important source for transmission of these organisms. First, hospitals are contaminated with clinically important multidrug-resistant organisms and *C difficile*.² Meticillin-resistant *Staphylococcus aureus* (MRSA), vancomycin-resistant enterococci (VRE), and *Acinetobacter* spp can survive on inanimate surfaces for days, and *C difficile* can survive for months.³ Second, only 50% of surfaces in hospital rooms are sufficiently cleaned between patient stays.⁴ As a result, patients admitted to rooms previously occupied by patients with multidrug-resistant organisms and *C difficile* are at an increased risk of subsequent infection or colonisation with these organisms.⁵ Finally, the contaminated environment is an important source of health-care personnel hand contamination.^{6–8}

Terminal room disinfection (disinfection of a room between occupying patients) can be enhanced by using a chemical disinfectant with sporicidal activity or by use of supplemental disinfection technologies. However, to our knowledge, no multicentre randomised assessment of enhanced terminal room disinfection strategies has been done.⁹ We designed the Benefits of Enhanced Terminal Room Disinfection study to assess the effects of four

different strategies for terminal room disinfection on acquisition of multidrug-resistant organisms and *C difficile*.

Methods

Study design and participants

We did this pragmatic, multicentre, cluster-randomised, crossover trial in nine hospitals in the USA from April, 2012, to July, 2014 (appendix). We tested one of four strategies for terminal room disinfection. Three strategies included enhanced terminal disinfection, and one included the standard terminal disinfection.

These four strategies were used in targeted rooms, defined as single-patient rooms from which a patient on contact precautions was discharged or transferred. In the reference group, targeted rooms were disinfected with quaternary ammonium-containing disinfectant for all rooms except those with patients with *C difficile*, in which a hypochlorite-containing disinfectant (bleach) was used. In the UV group, targeted rooms were disinfected with quaternary ammonium-containing disinfectant and a UV-C device except rooms of patients with *C difficile*, in which a bleach-containing disinfectant and UV-C device were used. In the bleach group, a bleach-containing disinfectant was used in all targeted rooms. In the bleach and UV group, a bleach-containing disinfectant and a UV-C device were used for all targeted rooms.

Each strategy was used at every study hospital for four consecutive 7-month study periods. Each study period consisted of a 1-month wash-in period followed by a 6-month period of data collection. The sequence of disinfection strategies was randomly selected for each hospital.

We selected study hospitals to include multiple types of hospitals (tertiary, community, Veterans Affairs) as a convenience sample. All microbiological cultures were considered for inclusion in our outcomes. Cultures may have been representative of infection or colonisation and included surveillance cultures, if obtained by policy at the study hospital. No screening cultures were obtained specifically for the study.

The Duke University Health System Institutional Review Board served as the central institutional review board. We received a waiver of informed consent for this study.

Randomisation and masking

We did resource-dependent randomisation of hospitals, taking into account the number of UV devices available (nine). First, we used a random number generator to determine the order in which hospitals would be randomly assigned a disinfection strategy. Then, we used a random number generator to determine the order in which disinfection strategies were used in each hospital. We continued this process for each hospital but counted the number of machines already assigned for other hospitals in each study period. If all nine UV-C devices were already assigned for a period, subsequent hospitals could not be assigned to one of the UV strategies for that period. Ultimately, all hospitals used all four strategies in a 1:1:1:1 ratio (appendix p 9). Allocation was not masked.

Procedures

All hospitals used gown and glove precautions (ie, contact precautions) for patients known or suspected to harbour multidrug-resistant organisms or *C difficile*. Environmental services personnel were trained on the appropriate use of the disinfectants, cleaning protocols, and UV-C device. The appendix provides information on standardisation of disinfection practices, implementation, and measures of protocol fidelity (appendix pp 1–2).

We did a microbiological analysis of 92 randomly selected seed rooms at two study hospitals to determine the total and average number of colony-forming units of the four target organisms that remained in the hospital room after terminal room disinfection (appendix p 4). Microbiological analyses and identification were done with standard protocols.¹⁰ All hospitals used PCR-based nucleic acid amplification tests to identify *C difficile* throughout the study.

We designed this study to detect infection or colonisation with one of four target organisms: MRSA, VRE, *C difficile*, or multidrug-resistant *Acinetobacter*.¹¹ A seed room was defined as a room containing a patient with microbiologically proven current or history of infection or colonisation with one or more target organisms. History of infection or colonisation was defined as any positive culture within the 12 months before admission. The next patient admitted to the seed room was an exposed patient. Community-onset was defined as the isolation of a target organism within the first 48 h of hospital admission. Hospital-acquired was defined as the isolation of a target organism after 48 h of hospital admission.

Outcomes

We had two primary outcomes: first, the incidence of all target organisms among patients exposed to seed rooms, and second, the incidence of *C difficile* infection among patients exposed to seed rooms, in the intention-to-treat population. Secondary outcomes were incidence among exposed patients of MRSA, of VRE, and of multidrug-resistant *Acinetobacter*; incidence in the whole hospital of all target organisms, of MRSA, of VRE, of *C difficile*, and of multidrug-resistant *Acinetobacter*; and adverse events (rate of UV-C device failure, time on diversion, emergency room wait time, health-care worker perception of cleaning methods, and room turnover time [time between patient discharge and completion of terminal room disinfection]; appendix pp 4–5). Incidence was calculated as the number of qualifying incident cases per 10 000 exposure days. Exposure days were calculated as the number of days the exposed patient spent in the seed room. Patients excluded from the numerator were also excluded from the denominator. Adverse outcomes were assessed at the hospital-level (ie, all patients or rooms were included in the analyses unless otherwise stated).

Three additional predetermined variables were measured at each study hospital: hand hygiene compliance, room cleaning compliance, and colonisation pressure (appendix pp 10–11).¹² We obtained demographic data and comorbid conditions for all exposed patients through administrative databases to calculate Charlson scores.¹³

We did two post-hoc analyses: (1) of the incidence of target organisms among exposed patients after removing the criteria requiring a minimum of 24 h in the seed room; and (2) of

the incidence of target organisms due to vegetative bacteria (MRSA, VRE, and multidrug-resistant *Acinetobacter*; appendix p 4).

Exposed patients qualified as an incident case of acquisition¹⁴ if they met the following criteria: in a seed room for 24 h or more AND a positive clinical culture or test with one of the target organisms AND the organism identified in the clinical culture or test was the same target organism isolated from the preceding patient in the seed room AND the positive culture or test was obtained during the index admission either during exposure to the seed room OR the positive culture or test was obtained after exposure to the seed room during the index admission or readmission within 90 days of discharge from the room for MRSA, VRE, and multidrug-resistant *Acinetobacter*¹⁵ or within 28 days of discharge from the room for *C difficile*.¹⁶ We excluded incident cases if they were community-onset infections or the exposed patient had a microbiologically proven history of infection or colonisation with the same target organism during the 12 months before admission.

Statistical analysis

We did power calculations based on a review of 4 years of surveillance data from study hospitals and published literature. All power calculations were done with two-sided significance level of 0.05. We projected that 1.96 million patient-days of care would be provided at the nine study hospitals (after excluding the wash-in periods). For each 6-month intervention period, we projected that approximately 491 200 patient-days of care would occur (distributed across nine participating hospitals). Based on data from our pre-existing surveillance databases, we projected that 959 outcomes due to the four target organisms would occur during the baseline (or reference) 6-month period (ie, with standard terminal room disinfection and no use of UV-C), for a baseline incidence of 1.95 per 1000 patient-days. Under these assumptions, the study would have 60% power to detect a 10% decrease in incidence rate, 92% power to detect a 15% decrease, and more than 99% power to detect a 20% decrease. The power analysis was done using simulation and was based on a Poisson regression model with hospital-level incidence rate as the outcome and disinfection strategy and hospital as the covariates.

All qualifying incident cases were included in the intention-to-treat population. The per-protocol population was identical to the intention-to-treat population for the reference strategy and bleach strategy. For the two strategies involving the UV device, the per-protocol population included qualifying incident cases who entered a seed room with documented use of the UV device. For the purposes of this study, the UV device only had to be turned on, the cycle did not have to be completed. The appendix contains a more detailed discussion of the differences between the analysis populations (pp 2–3).

We summarised patient characteristics using percentages for categorical variables and medians for continuous variables. We analysed outcomes using intention-to-treat and per-protocol principles for outcomes among exposed patients. We analysed incidence rates using overdispersed Poisson models with disinfection group (reference, UV, bleach, and bleach and UV), order of the strategies within the study (whether a particular strategy was used in the first, second, third, or fourth study period), and hospital as fixed-effect categorical covariates. We used generalised estimating equations to account for correlation between

different study strategies within the same hospital. Each of the study groups was compared to the reference group except for the analysis of *C difficile* among exposed patients. Because this comparison involved the comparison of bleach vs bleach and UV-C, results from the UV group and the bleach and UV group were compared to results from the reference group and the bleach group. We used the same model construction strategy for all outcome analyses. We calculated relative risk (RR), 95% CIs, and risk reductions for each model. Statistical tests were done at a two-sided significance level of 0.05. In light of the pragmatic nature of the trial, we made no adjustments for multiple comparisons. We did all statistical analyses using SAS (version 9.4).

The study is registered on ClinicalTrials.gov (NCT01579370).

Role of the funding source

The funder served an advisory role in the development of the study protocol. All authors had full access to all data in the study and had final responsibility for the decision to submit for publication.

Results

All nine hospitals participated in the study from April, 2012, to July, 2014. The appendix shows the randomised assignment for each hospital (p 13). The average cluster size was 7807 (SD 644) patients. 31 226 patients were exposed to a seed room; 24 585 (79%) stayed in the seed room for 24 h or more, and 21 395 (69%) met all inclusion criteria (figure). Baseline characteristics of qualifying exposed patients were similar for all four cleaning strategies (table 1).

A total of 423 outcomes were recorded: 228 (54%) cultures represented infection and 195 (46%) represented colonisation. 115 patients had a primary outcome during 22 426 exposure days during the reference period (51.3 per 10 000 exposure days); the median incidence of target organisms in the baseline period per hospital was 37.1 per 10 000 exposure days (range 17.5–101.6). The addition of a UV-C device to the standard disinfection strategy significantly decreased the incidence of target organisms to 33.9 per 10 000 exposure-days (n=76; RR 0.70, 95% CI 0.50–0.98; p=0.036; table 2; appendix p 14). The incidence of target organisms was lower in eight of the nine study hospitals in the UV group (appendix p 15).

There was no significant difference in the incidence of target organisms from rooms treated with bleach compared with reference (table 2). Similarly, there was no significant difference between use of bleach and UV compared with reference (table 2). The appendix shows outcomes from individual study hospitals by intention to treat for each disinfection strategy (p 7).

The incidence of *C difficile* was not significantly different with or without UV-C devices (table 2). The incidence of MRSA was not significantly lower in the UV group and essentially unchanged in the bleach and bleach and UV groups (table 2). The incidence of VRE was not significantly lower in the UV group but was significantly lower in both groups

that used a bleach-containing disinfectant (table 2). The use of bleach decreased the incidence of VRE by 57% compared to reference; the use of bleach and a UV-C device decreased incidence of VRE by 64% (table 2). Only one patient acquired multidrug-resistant *Acinetobacter* after exposure in a seed room. Thus, no comparisons or models were constructed for this organism.

2848 (55%) of 5178 eligible rooms in the UV group and 3701 (63%) of 5863 eligible rooms in the bleach and UV group were included in the per-protocol analyses. Effect estimates were generally similar in per-protocol analyses to the intention-to-treat analyses (table 3). The incidence of MRSA, however, was significantly lower in the UV group compared with the reference group (table 3).

Our microbiological assessment of 92 seed rooms after terminal disinfection showed that all enhanced strategies decreased the bioburden of target organisms, but the largest decrease occurred in the UV group (table 4). Protocol compliance, hand hygiene compliance, cleaning compliance, and colonisation pressure were similar across study groups (table 5; appendix p 4).

The median room cleaning time was approximately 4 min longer in the UV and UV and bleach groups (table 5). The total wait time in the emergency department and days on diversion were unchanged across disinfection strategies. Time from admit decision to departure from the emergency department was approximately 10–20 min longer in each of the enhanced disinfection groups compared with the reference group. One hospital reported a single UV-C exposure event during the study (appendix p 5). Additional secondary analyses, including incidence in the whole hospital of all target organisms, of MRSA, of VRE, of *C difficile*, and of multidrug-resistant *Acinetobacter*, and health-care worker perception of cleaning methods, will be presented elsewhere.

We did two post-hoc analyses (appendix p 9). First, removing the 24-h exposure requirement for exposed patients did not change the effect measures. Second, after excluding patients admitted to *C difficile* seed rooms, the decrease in incidence of target vegetative multidrug-resistant organisms was strengthened in the UV group and significantly lower in the bleach and UV group.

Discussion

Our large, prospective, multicentre, cluster-randomised trial is the first, to our knowledge, to demonstrate a decrease in acquisition and infection with epidemiologically important pathogens following the use of enhanced room disinfection strategies. Patients admitted to rooms previously occupied by patients harbouring a multidrug-resistant organism or *C difficile* were 10–30% less likely to acquire the same organism if the room was terminally disinfected using an enhanced strategy. The largest risk reduction occurred when a UV-C device was added to the standard disinfectant strategy. By contrast, we showed no statistically significant decrease in outcomes when we used enhanced terminal disinfection with bleach or bleach and UV. Similarly, the incidence of *C difficile* infection was not different among exposed patients after adding UV to bleach disinfection.

Our results need to be interpreted in the appropriate context. First, decreases in acquisition of target organisms associated with the use of enhanced disinfection strategies were recorded even though our reference group was also an enhanced strategy of sorts. Overall compliance with thoroughness of cleaning in the reference group was roughly 90%. By contrast, most previous studies conclude that approximately half of all hospital room surfaces are not cleaned during terminal cleaning.⁴ Improved cleaning compliance decreases environmental bioburden¹⁷ and risk of acquisition, particularly of MRSA and VRE.^{18,19} Second, in the reference group, the quaternary ammonium-containing disinfectant was delivered with microfibre cloths, which remove more bacteria than cotton and synthetic fibre cloths.²⁰ Third, the enhanced nature of the reference group and lack of multidrug-resistant *Acinetobacter* outcomes probably led to a decrease in power. Thus, the absence of a decrease in the incidence of target organisms among exposed patients in the bleach and bleach and UV groups might have been related to type II error.

No randomised controlled trials have previously been done using a UV device or enhanced chemical disinfectant. To our knowledge, only one other randomised controlled trial has investigated an enhanced terminal room disinfection strategy. A hydrogen peroxide vapour system was evaluated over 30 months in six high-risk units in a single tertiary care centre.²¹ Patients in intervention units had a 64% decrease in acquisition of multidrug-resistant organisms and *C difficile* and a 75% decrease in acquisition of VRE compared to patients in control units.

UV devices reduce the environmental bioburden of MRSA, VRE, *C difficile*, and *Acinetobacter* spp.^{10,22} Of four published studies on the clinical effectiveness of UV devices, one showed a 20% decrease in hospital-acquired multidrug-resistant organisms²³ and three showed 22–53% decreases in *C difficile* infection.^{24–26} In light of these results, we were surprised by the lack of change in rates of *C difficile* among exposed patients. This lack of change might have been caused by the following factors. First, the reference group for our *C difficile*-specific outcome involved the use of bleach. As we had high (around 90%) compliance with the use of bleach, there may have been relatively few residual spores for the UV device to eliminate.¹⁷ Second, UV is less effective against *C difficile* than against vegetative bacteria, especially in areas of shadow.^{10,22} Third, we used a single-stage cycle with the UV-C device placed adjacent to but outside of the bathroom.²⁷ Thus, we may not have effectively eliminated *C difficile* from bathrooms. Finally, the environment might not play as large a role in *C difficile* transmission as previously suspected.²⁸ Eyre and colleagues²⁹ assessed 1250 cases of symptomatic *C difficile* in Oxfordshire, UK, over a 4-year period using whole genome sequencing and reported that 45% of *C difficile* cases were genetically distinct from previous cases. Although this analysis did not consider asymptomatic colonisation, only 2% of patients with related *C difficile* isolates were linked by possible environmental contamination. Our post-hoc analysis excluding patients exposed to *C difficile* showed that the effect in the UV group was strengthened and the effect in the bleach and UV group became statistically significant.

To our knowledge, no other randomised controlled trials have assessed the effect of using a sporicidal disinfectant on the incidence of our target organisms. Grabsch and colleagues³⁰ recorded a 67% decrease in acquisition of VRE and an 83% decrease in VRE bacteraemia

with use of bleach. Results from the intention-to-treat analysis in our trial did not show a significant decrease in the incidence of target organisms following the routine use of bleach for terminal disinfection of contact precaution rooms. Results from a prespecified secondary analysis, however, validate the decrease in VRE reported by Grabsch and colleagues. Our clinical results were largely corroborated by our microbiological assessment of seed rooms after terminal disinfection; greater reductions in colony-forming units occurred in the UV group than in the bleach and UV and bleach groups.

Our study has limitations. First, we relied on clinical cultures obtained during the course of standard care, which might have introduced ascertainment bias. Clinicians might have changed their culturing practices during the course of the study. Additionally, because we did not screen seed patients with a history of infection or colonisation or all exposed patients on exit from the seed rooms, we did not capture all acquisitions, might have failed to exclude a patient with community-onset colonisation, and our denominators might have included extra exposure days. We doubt, however, that any of these scenarios affected our results given randomisation. Second, we did not do molecular analyses to confirm that the organisms included in our outcomes were related to organisms in the environment, as this task was impossible given the scope of our study. Third, we did not account for multiplicity in our statistical testing given the pragmatic nature of our study; thus, the p values generated from our analyses should be interpreted with caution. Fourth, as noted above, our study had an enhanced reference group and thus decreased power. As a result, we suspect the effect measures in our study represent minimum effects of these strategies. Finally, our intervention was directed towards three multidrug-resistant organisms and *C difficile*. We suspect that enhanced terminal room disinfection strategies decrease risk of acquisition of non-multidrug-resistant organisms, such as methicillin-susceptible *S aureus* and vancomycin-susceptible enterococci.

Acquisition and infection with multidrug-resistant organisms and *C difficile* in health care is a complex and multifaceted process. Our study suggests that (1) the health-care environment is an important source for acquisition of multidrug-resistant organisms and *C difficile*, and (2) the risk of acquisition of these pathogens from the environment can be modified. More than a century after Semmelweis and Lister's landmark studies, our results suggest that methods to improve disinfection can still lead to better patient outcomes.

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Research in context

Evidence before this study

The hospital environment is contaminated with multidrug-resistant organisms and *Clostridium difficile* and is often disinfected inadequately. As a result, patients who enter contaminated hospital rooms are at increased risk for acquisition and infection from these organisms. Enhanced disinfection strategies may decrease the risk for transmission of such bacteria through the hospital environment, but supportive evidence is limited to single centre or quasi-experimental studies. According to a systematic review by Han and colleagues, no randomised multicentre trials have been done to determine the efficacy of enhanced strategies.

Added value of this study

Our study is, to our knowledge, the first multicentre randomised controlled trial to evaluate the effect of enhanced disinfection strategies on acquisition and infection due to four target organisms (meticillin-resistant *Staphylococcus aureus*, vancomycin-resistant staphylococci, multidrug-resistant *Acinetobacter*, and *C difficile*). Adding a UV-C device to quaternary ammonium disinfection decreased the risk of subsequent acquisition and infection by target organisms. Our study shows the efficacy of enhanced disinfection and confirms that the contaminated hospital environment is a modifiable risk factor.

Implications of all the available evidence

Multidrug-resistant organisms and *C difficile* lead to adverse patient outcomes. Novel and improved prevention strategies are needed. Prevention of the spread of these organisms will probably require a multifaceted approach, including enhanced disinfection, improved hand hygiene, and antimicrobial stewardship.

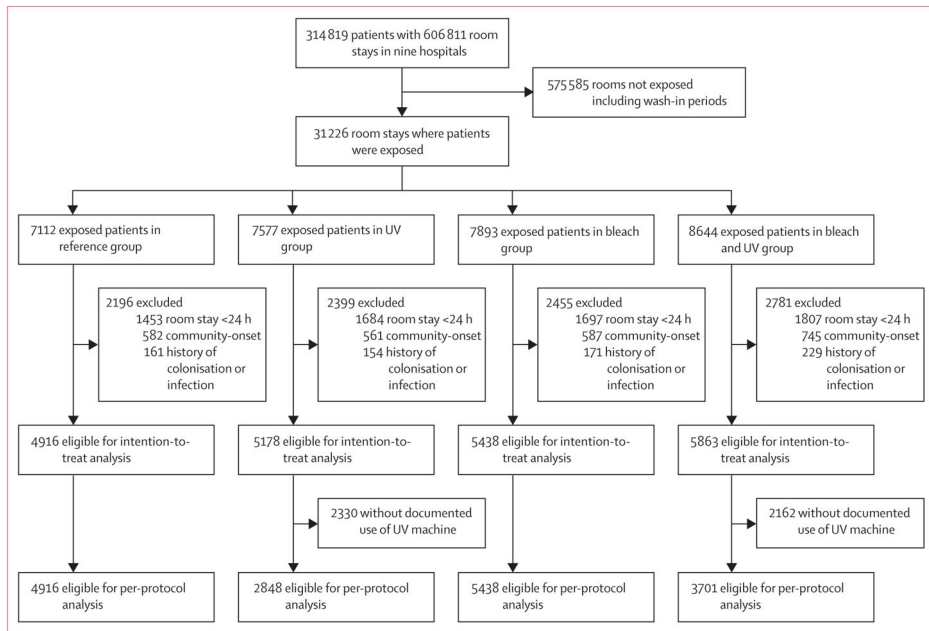


Figure.
Trial profile

Table 1

Baseline characteristics

	Reference (n=4916)	UV group (n=5178)	Bleach group (n=5438)	Bleach and UV group (n=5863)
Demographics*				
Mean age (SD)	57.9 (20.9)	58.5 (21.3)	58.6 (20.7)	57.7 (21.8)
Race				
White	3042 (63%)	3228 (65%)	3416 (64%)	3747 (64%)
African American	1418 (30%)	1411 (28%)	1591 (30%)	1655 (28%)
Other	243 (5%)	233 (5%)	249 (5%)	329 (6%)
Unknown	102 (2%)	97 (2%)	95 (2%)	111 (2%)
Male sex	2475 (51%)	2518 (51%)	2768 (52%)	3017 (52%)
Comorbidities*				
Median Charlson index (IQR)	2 (1–4)	2 (0–4)	2 (1–4)	2 (0–4)
Myocardial infarction	499 (11%)	457 (10%)	475 (9%)	583 (10%)
Congestive heart failure	937 (20%)	950 (20%)	1014 (20%)	1151 (21%)
Cerebrovascular disease	571 (12%)	540 (11%)	582 (11%)	610 (11%)
Hemiplegia or paraplegia	97 (2%)	118 (2%)	139 (3%)	166 (3%)
Peripheral vascular disease	450 (10%)	498 (10%)	524 (10%)	543 (10%)
Dementia	75 (2%)	101 (2%)	138 (3%)	111 (2%)
COPD	1248 (27%)	1325 (28%)	1339 (26%)	1516 (27%)
Rheumatic disease	161 (3%)	181 (4%)	183 (4%)	224 (4%)
Peptic ulcer disease	143 (3%)	97 (2%)	126 (2%)	178 (3%)
Liver disease				
Mild	475 (10%)	452 (9%)	484 (9%)	557 (10%)
Moderate or severe	120 (3%)	142 (3%)	135 (3%)	177 (3%)
Diabetes mellitus	1302 (28%)	1248 (26%)	1371 (27%)	1505 (27%)
Complicated	303 (7%)	273 (6%)	350 (7%)	350 (6%)
Renal disease	980 (21%)	986 (21%)	1083 (21%)	1171 (21%)
Malignancy	842 (18%)	807 (17%)	864 (17%)	961 (17%)
Metastatic solid tumour	305 (7%)	340 (7%)	330 (6%)	367 (7%)
AIDS/HIV	49 (1%)	48 (1%)	52 (1%)	55 (1%)
Arrhythmia	1542 (33%)	1425 (30%)	1513 (30%)	1824 (33%)
Valvular heart disease	420 (9%)	415 (9%)	412 (8%)	566 (10%)
Pulmonary circulation disease	386 (8%)	452 (9%)	439 (9%)	495 (9%)
Hypertension	2264 (49%)	2237 (47%)	2336 (46%)	2709 (48%)
Complicated	826 (18%)	839 (18%)	959 (19%)	1007 (18%)
Other neurological disease	682 (15%)	665 (14%)	671 (13%)	792 (14%)
Hypothyroid disease	523 (11%)	545 (11%)	599 (12%)	637 (11%)

Data are n (%) unless stated otherwise.

* The majority of patient-specific data were not available from one study hospital because of changes in electronic health record systems as follows: age (406 had data missing), race (428 missing), sex (407 missing), comorbidity data (1267 missing).

COPD=chronic obstructive pulmonary disease.

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Table 2

Results of intention-to-treat analysis

	Reference	UV group	Bleach group	Bleach and UV group
All target organisms				
Exposed patients	4916	5178	5438	5863
Incident cases (%)	115 (2.3%)	76 (1.5%)	101 (1.9%)	131 (2.2%)
Exposure days	22 426	22 389	24 261	28 757
Rate (per 10 000 exposure-days)	51.3	33.9	41.6	45.6
Risk reduction (95% CI)	Reference	17.4 (5.8 to 28.9)	9.7 (-2.7 to 22.0)	5.7 (-6.2 to 17.7)
RR (95% CI); p value	Reference	0.70 (0.50 to 0.98); 0.036	0.85 (0.69 to 1.04); 0.116	0.91 (0.76 to 1.09); 0.303
<i>Clostridium difficile</i> *				
Exposed patients	2499	2678
Incident cases (%)	36 (1.4%)	38 (1.4%)
Exposure days	11 385	12 509
Rate (per 10 000 exposure-days)	31.6	30.4
Risk reduction (95% CI)	Reference	1.2 (-12.7 to 15.2)
RR (95% CI); p value	Reference	1.0 (0.57 to 1.75); 0.997
Meticillin-resistant <i>Staphylococcus aureus</i>				
Exposed patients	3300	3451	3631	3848
Incident cases (%)	73 (2.2%)	54 (1.6)	74 (2.0)	89 (2.3)
Exposure days	14 524	14 780	15 343	18 960
Rate (per 10 000 exposure-days)	50.3	36.5	48.2	46.9
Risk reduction (95% CI)	Reference	13.8 (0.1 to 27.3)	2.1 (-13.8 to 17.8)	3.4 (-8.9 to 15.5)
RR (95% CI); p value	Reference	0.78 (0.58 to 1.05); 0.104	1.00 (0.82 to 1.21); 0.967	0.97 (0.78 to 1.22); 0.819
Vancomycin-resistant enterococci				
Exposed patients	1055	1206	1468	1753
Incident cases (%)	37 (3.5%)	17 (1.4%)	24 (1.6%)	37 (2.1%)
Exposure days	5838	5780	7522	9488
Rate (per 10 000 exposure-days)	63.4	29.4	31.9	39.0
Risk reduction (95% CI)	Reference	34.0 (9.3 to 58.6)	31.5 (12.7 to 50.2)	24.4 (0.5 to 48.2)
RR (95% CI); p-value	Reference	0.41 (0.15 to 1.13); 0.084	0.43 (0.19 to 1.00); 0.049	0.36 (0.18 to 0.70); 0.003
Multidrug-resistant <i>Acinetobacter</i> †				
Exposed patients	31	47	28	62
Incident cases (%)	0	0	1 (3.6)	0
Exposure days	156	199	98	244
Rate (per 10 000 exposure-days)	0	0	102.4	0

RR=relative risk.

* Rooms with patients known or suspected of having *C difficile* infection were terminally cleaned with bleach-containing solutions in all study disinfection strategies.

† We created no models for multidrug-resistant *Acinetobacter* because only one outcome occurred in the nine study hospitals across all four study groups.

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Table 3

Results of per-protocol analysis

	Reference group	UV group	Bleach group	Bleach and UV group
All target organisms				
Exposed patients	4916	2848	5438	3701
Incident cases (%)	115 (2.3%)	46 (1.6%)	101 (1.9%)	93 (2.5%)
Exposure days	22 426	12 299	24 261	17 354
Rate (per 10 000 exposure-days)	51.3	37.4	41.6	53.6
Risk reduction (95% CI)	Reference	13.9 (−0.1 to 27.9)	9.7 (−2.7 to 22.0)	−2.3 (−15.7 to 11.1)
RR (95% CI); p value	Reference	0.69 (0.50 to 0.95); 0.025	0.74 (0.61 to 0.91); 0.004	1.0 (0.81 to 1.23); 1.00
<i>Clostridium difficile</i> *				
Exposed patients	2499	1712
Incident cases (%)	36 (1.4%)	30 (1.8%)
Exposure days	11 385	8015
Rate (per 10 000 exposure-days)	31.6	37.4
Risk reduction (95% CI)	Reference	−5.8 (−17.1 to 5.5)
RR (95% CI); p value	Reference	1.22 (0.68 to 2.17); 0.511
Meticillin-resistant <i>Staphylococcus aureus</i>				
Exposed patients	3300	1872	3631	2425
Incident cases (%)	73 (2.2%)	28 (1.5%)	74 (2.0%)	63 (2.6%)
Exposure days	14 525	7934	15 343	10 681
Rate (per 10 000 exposure-days)	50.3	35.3	48.2	59.0
Risk reduction (95% CI)	Reference	15.0 (−0.6 to 30.6)	2.1 (−13.8 to 17.8)	−8.7 (−18.0 to 0.5)
RR (95% CI); p value	Reference	0.67 (0.48 to 0.94); 0.019	0.89 (0.72 to 1.09); 0.260	1.09 (0.85 to 1.39); 0.503
Vancomycin-resistant enterococci				
Exposed patients	1055	659	1468	1134
Incident cases (%)	37 (3.5%)	13 (2.0%)	24 (1.6%)	24 (2.1%)
Exposure days	5838	3265	7522	6237
Rate (per 10 000 exposure-days)	63.4	39.8	31.9	38.5
Risk reduction (95% CI)	Reference	23.6 (−6.1 to 53.2)	31.5 (12.7 to 50.2)	24.9 (−0.6 to 50.4)
RR (95% CI); p value	Reference	0.56 (0.21 to 1.50); 0.248	0.35 (0.16 to 0.78); 0.010	0.41 (0.22 to 0.77); 0.006

Data are unchanged for multidrug-resistant *Acinetobacter baumannii* (table 2).

* Rooms with patients known or suspected of having *C difficile* infection were terminally cleaned with hypochlorite-containing solutions.

Table 4

Microbiological assessment

	Reference (n=21)		UV group (n=28)		Bleach group (n=23)		Bleach and UV group (n=20)	
	Total CFU	Mean CFU per room (SD)	Total CFU	Mean CFU per room (SD)	Total CFU	Mean CFU per room (SD)	Total CFU	Mean CFU per room (SD)
<i>Clostridium difficile</i>	79	3.8 (14.2)	80	2.9 (12.6)	103	4.5 (14.5)	65	3.3 (10.4)
MRSA	179	8.5 (27.1)	3	0.1 (0.6)	101	4.4 (15.0)	17	0.9 (2.5)
VRE	831	39.6 (127.5)	6	0.2 (0.8)	56	2.4 (5.6)	38	1.9 (6.1)
MDR <i>Acinetobacter</i> spp	188	9.0 (36.4)	5	0.2 (0.9)	9	0.4 (1.9)	5	0.3 (0.9)
Total target organisms	1277	60.8 (161.3)	94	3.4 (13.4)	269	11.7 (21.4)	125	6.3 (16.1)

MRSA=meticillin-resistant *Staphylococcus aureus*. VRE=vancomycin-resistant enterococci. MDR=multidrug-resistant. CFU=colony-forming units.

Table 5

Hospital outcomes

	Reference	UV group	Bleach group	Bleach and UV group
Hospital-level variables				
Hand hygiene compliance				
Observations	59 519	64 810	64 950	57 650
Median per hospital (IQR)	89.7 (86.3–94.7)	88.1 (84.6–95.7)	91.4 (86.2–96.1)	90.8 (82.8–94.1)
Room cleaning				
Room observations	5717	4312	4538	5869
Mean number of locations monitored per room	12.4	11.6	12.5	12.0
Locations monitored	70 704	50 081	56 753	70 312
Median compliance (IQR)	100% (91–100)	95% (86–100)	100% (87–100)	100% (84–100)
Median colonisation pressure (IQR)	4.6% (3.1–9.7)	4.3% (3.6–6.6)	4.5% (3.7–5.8)	4.8% (3.4–6.7)
Protocol compliance				
pH pen use (%)	4836/5024 (96%)	2970/3262 (91%)	1161/1206 (96%)	5899/6002 (98%)
Median (IQR)	100 (91–100)	93 (89–97)	96 (95–98)	99 (97–99)
UV-C devices used in contact precaution rooms (%)	..	6214/7137 (87%)	..	10 006/11 274 (89%)
Median (IQR)	..	92 (85–93)	..	91 (87–91)
Adverse events				
Room turnover times				
Median total turnover time (IQR)	79.4 (74.4–117.3)	88.9 (80.0–93.4)	82.6 (73.1–123.3)	87.5 (76.2–127.0)
Rooms	78 413	127 028	114 101	102 227
Median room cleaning time (IQR)	35.9 (32.5–38.5)	40.7 (38.4–42.1)	35.6 (32.3–38.9)	40.1 (39.1–44.2)
Rooms	133 744	144 183	132 753	137 814
Emergency department waiting times (min)				
Median total time in emergency department (range; n=7)	392 (290–537)	390 (286–534)	392 (290–533)	399 (294–544)
Observations	34 532	31 961	30 613	32 320
Median time from admit decision to departure from emergency department (range; n=4)	92 (64–135)	110 (72–180)	116 (75–194)	108 (70–184)
Observations	18 443	24 025	21 566	23 732
Time on diversion (days)				
Total	63.8	53.8	34.2	38.2
Median per hospital (IQR)	2.7 (0.3–11.7)	2.5 (1.7–7)	2.9 (0.6–7.5)	1.3 (0.6–6.6)