Analytical Performance Issues
Comparison of ATP Bioluminescence and Aerobic Bacterial Count for Evaluating Surface Cleanliness in an Italian Hospital

Contaminated hospital surfaces have been demonstrated to be an important environmental reservoir of microorganisms that can increase the risk of nosocomial infection in exposed patients. As a consequence, cleaning and disinfecting hospital environments play an important role among strategies for preventing healthcare-associated colonization and infections. The aim of the present study was to evaluate whether adenosine triphosphate (ATP) presence, measured by bioluminescence methods, can predict microbiological contamination of hospital surfaces. The study was carried out between September and December 2012 at the University Hospital “P. Giaccone” of Palermo.

A total of 193 randomly selected surfaces (tables, lockers, furnishings) were sampled and analyzed in order to assess ATP levels (expressed as relative light units or RLU) and aerobic colony count (ACC) or presence of *S. aureus*. ACC had median values of 1.85 cfu/cm² (interquartile range = 4.16) whereas ATP median was 44.6 RLU/cm² (interquartile range = 92.3). Overall, 85 (44.0%) surfaces exceeded the established microbial benchmark: 73 (37.8%) exceeded the 2.5 cfu/cm² ACC standard, 5 (2.6%) surfaces were positive for *S. aureus* and 7 (3.6%) showed both the presence of *S. aureus* and an ACC of more than 2.5 cfu/cm². ACC and bioluminescence showed significant differences in the different surface sites (*p* < 0.001). A significant correlation was found between ACC and RLU values (*p*-value < 0.001; *R*² = 0.29) and increasing RLU values were significantly associated with a higher risk of failing the benchmark (*p* < 0.001).

Our data suggest that bioluminescence could help in measuring hygienic quality of hospital surfaces using a quick and sensitive test that can be an useful proxy of microbial contamination; however, further analysis will be necessary to assess the cost-efficacy of this methodology before requiring incorporation in hospital procedures.

INTRODUCTION

Contaminated hospital surfaces have been demonstrated to be an important environmental reservoir of microorganisms that can contribute to increase the risk of nosocomial infection in exposed patients.\(^{(1)}\) As a consequence, cleaning and disinfecting hospital environments play an important role among strategies for preventing healthcare-associated colonization and infections.\(^{(2)}\) A major problem in defining these recommendations is to classify when a surface can be considered “clean” and “acceptable.” Although, a large number of international guidelines provide for visual surface inspection for measuring cleanliness,\(^{(3)}\) this method cannot be considered an accurate measure of surface cleanliness or of microbial contamination and previous studies have indicated that it may overestimate cleaning efficacy,\(^{(4)}\) failing to predict the risk of infection for patients.\(^{(5)}\) Several other tools and methods including microbiological methods (e.g., aerobic bacterial counts, culture of indicator organisms) and bioluminescence tests for assessing cleanliness on a more scientific basis have been proposed. Microbiological cultures are commonly thought to

---

Column Editor
Martin Harper

Reported by
Emanuele Amodio\(^1\)
Lucia Cannova\(^1,2\)
Maria Rosaria Villafrate\(^1,2\)
Anna Maria Merendino\(^1,2\)
Luigi Aprea\(^1\)
Giuseppe Calamusa\(^1,2\)

\(^1\)Department of Sciences for Health Promotion, “G. D’Alessandro”, University of Palermo, Palermo, Italy

\(^2\)Sanitary Management of the Azienda Ospedaliera Universitaria, “P. Giaccone” of Palermo, Palermo, Italy

Address correspondence to: Emanuele Amodio, Department of Sciences for Health Promotion “G. D’Alessandro,” University of Palermo, Italy, Via Del Vespro n°133, cap 90127; e-mail: emanuele.amodio@unipa.it
adequately describe the infectious risk attributable to contaminated surfaces but they are time-consuming, needing a minimum of 24 hours.

Otherwise, bioluminescence tests, assessing the adenosine triphosphate (ATP) presence, provide quick and objective feedback on surface cleanliness, reporting the presence of organic materials including microbiological contamination. Few studies have evaluated ATP bioluminescence methods for monitoring cleanliness in hospitals but the relationship between bioluminescence and microbiological results remains poorly characterized.\(^\text{4,6–8}\) The purpose of the present study was to evaluate whether ATP presence, measured by bioluminescence methods, can predict microbial contamination of hospital surfaces.

**METHODS**

**Study Design**

The study was carried out between September and December 2012 at the University Hospital “P. Giaccone” of Palermo that includes 72 hospital Units accounting for over 500 beds.

Surfaces to sample were randomly selected proportionally to the size of the each Unit. Sampling was performed during the morning (7:00 a.m. to 11:00 a.m.) about 2 hours after sanitization. Surface sites included tables, lockers, and furnishings (e.g., bed, chairs) in patient and healthcare workers’ rooms. Each site was sampled in order to measure aerobic colony count (ACC), ATP, and *Staphylococcus aureus* presence.

**Bioluminescence and Microbiological Analyses**

ATP levels were measured using Lumincontrol II (PBI International, Milano, Italy) and expressed as relative light units (RLU). A swab for ATP revelation in a 10×10 cm surface sampled by a close zig-zag pattern according to the manufacturer’s guidelines.

Microbiological assessment was based upon growth on slides pre-coated by the manufacturer (OXOID, Cambridge, UK) with Plate count agar and Mannitol Salt Agar with addition of germicide inhibitors. Each slide (55 mm) was pressed for 15 seconds onto the surface adjacent to that where ATP was determined and then incubated aerobically at 37°C for 48 hours. Colony density was determined by visual count according to the manufacturer’s standards. Each colony present in Mannitol Salt Agar and suggestive for *S. aureus* was then isolated in a diagnostic sensitivity test Agar and then identified and confirmed by detection of bound coagulase/protein A and a biochemical specific test for bacterial identification (Api Staph-BIOMERIEUX, Florence, Italy).

Surfaces were considered at risk for determining infection when they had an ACC of at least 2.5 cfu/cm\(^2\) or presence of *S. aureus* as an indicator organism of pathogen contamination.\(^\text{5,9–11}\)

**Statistical Analyses**

Data were analyzed using the R statistical software package.\(^\text{12}\) Statistical significance was defined as \(p \leq 0.05\), two-tailed. Absolute and relative frequencies were calculated for qualitative variables. Quantitative variables were summarized as median (interquartile range). Frequencies were analyzed by Chi-square test whereas medians were compared by using the Mann-Whitney Rank sum test. Pearson correlation coefficient and linear regression model were used to describe the relationship between logarithmic transformed ACC and ATP values. Finally, ACC > 2.5 cfu/cm\(^2\) or presence of *S. aureus* were considered as S standard microbial benchmark indicating significant environmental contamination.

**RESULTS**

During the study period, a total of 193 surfaces were analyzed: 89 tables (46.1%), 50 lockers (25.9%), and 54 furnishings (28%). ACC had median values of 1.85 cfu/cm\(^2\) (interquartile range = 4.16) whereas ATP median was 44.6 RLU/cm\(^2\) (interquartile range = 92.3).

Overall, 85 (44.0%) surfaces exceeded the established microbial benchmark: 73 (37.8%) exceeded the 2.5 cfu/cm\(^2\) ACC standard, 5 (2.6%) surfaces were positive for *S. aureus* and 7 (3.6%) showed both the presence of *S. aureus* and an ACC of more than 2.5 cfu/cm\(^2\). Although surfaces contaminated with *S. aureus* showed higher ACC and ATP medians (3.5 vs. 2 cfu/cm\(^2\) and 51.3 vs. 43.5 RLU/cm\(^2\), respectively), these associations were not statistically significant (\(p = 0.057\) and \(p = 0.36\), respectively).

As shown in Figure 1, both ACC and bioluminescence showed significant differences in the three surface sites (\(p < 0.001\) in both cases). In particular, lockers and tables were more contaminated with microbiological (median ACC = 5.5 and 2.5 cfu/cm\(^2\), respectively) and organic (median ATP = 61.3 and 46.1 RLU/cm\(^2\), respectively) materials.

Figure 2 shows the correlation between ACC and logarithmic transformed RLU values (F-statistic=80.3; p-value < 0.001, with a R\(^2\) value of 0.29). Figure 3 presents the distribution of samples that failed the established a microbial benchmark (ACC > 2.5 cfu/cm\(^2\) or presence of *S. aureus*) in relation to their ATP values.

Increasing RLU values were significantly associated with a higher risk of failing the benchmark (\(p < 0.001\)). The totality of samples with RLU values <100 passed the benchmark whereas more than 40% of samples with RLU >1,000 failed the standard.

**DISCUSSION**

Although hygiene of hospital surfaces is advocated worldwide as necessary to control nosocomial infections, to date there are still several concerns about which methods are preferred in assessing environmental cleanliness. As previously reported, visual assessment is probably the less accurate measure of surface cleanliness\(^\text{4,5}\) whereas microbiological assessments could be considered a reference method, being able to predict, at least theoretically, the risk of infection for patients. In the last decades several authors have...
proposed detection of ATP bioluminescence for quickly monitoring and having a standardized sensitive measure of hospital cleanliness. However, among the few studies that compared this method with microbiological cultures, a large majority found no correlation between RLU values and ACC.\(^{13-15}\) Otherwise, a minority of authors reported a good agreement or a weak relationship between the two methods.\(^{16,17}\) Our results confirm the latter findings, suggesting that surfaces with low microbial contamination also had low bioluminescence and that RLU values could explain about one-third of the ACC.
FIGURE 3. Distribution of samples that failed or passed the established microbial benchmark (ACC $>2.5$ cfu/cm² or presence of $S$. aureus) stratified for different ATP values.

variability. According to this observation, Griffith et al. found that about 33% of the ATP from hand contact surfaces is likely to be of microbial origin with the remainder non-microbial.\textsuperscript{(4)}

The fact that a part of measured bioluminescence is usually due to microbial presence could be useful in setting a minimum ATP value suggesting low microbial contamination. Furthermore, benchmark values of RLU 100,\textsuperscript{(17)} 250,\textsuperscript{(9)} and 500 RLU\textsuperscript{(3,4,14)} for ATP testing have been proposed to identify surfaces with a significant bacterial contamination, as recommended by international guidelines. The present study shows that a certain number of surfaces with ATP values $\leq 500$ RLU could have a microbial contamination higher than 2.5 cfu/cm² or presence of $S$. aureus and, consequently, the use of such a benchmark could be questionable. Otherwise, our experiences suggest that, using Lumicontrol II, an ATP value of 100 RLU could be a better bioluminescence benchmark, having a low risk ($<5\%$) of classifying as “clean” a significantly contaminated surface.

All these considerations could be affected by some limitations. First, the culture plates used are a selective filter and only indicate presence of organisms that may be culturable on such a media whereas no information is provided about those that are not culturable (e.g., anaerobic organisms and viable but non-culturable organisms). Second, our study has not considered standardized experimental contamination but, instead, included surfaces cleaned by various personnel and highly contaminated by patients, personnel, and visitors. This last point can also be considered a strength since it describes the real surface situation in a hospital environment.

CONCLUSION

Our data suggest that bioluminescence could help in measuring the hygienic quality of hospital surfaces using a quick and sensitive test that can be an useful proxy of microbial contamination; however, further analysis will be necessary to assess the cost-efficacy of this methodology before requiring its incorporation in hospital procedures.

ACKNOWLEDGMENTS

Thanks are due to Dr. Salvatore Casalicchio and Dr. Dario Cottitto for their helpful contributions to the study.

REFERENCES

1. Boyce, J.M.: Environmental contamination makes an important contribution to hospital infection. \textit{J. Hosp. Infect.} 65:50–54 (2007).
2. Hota, B.: Contamination, disinfection, and cross-colonisation? Are hospital surfaces reservoirs for nosocomial infection? \textit{Clin. Infect. Dis.} 39:1182–1189 (2004).
3. Cooper, R.A., C.J. Griffith, R.E. Malik, P. Obbe, and N. Looker: Monitoring the effectiveness of cleaning in four British hospitals. \textit{Am. J. Infect. Control} 35(5):338–341 (2007).
4. Griffith, C.J., R.A. Cooper, J. Gilmore, C. Davies, and M. Lewis: An evaluation of hospital cleaning regimes and standards. \textit{J. Hosp. Infect.} 45:19–28 (2000).
5. Dancer, S.J.: How do we assess hospital cleaning? A proposal for microbiological standards for surface hygiene in hospitals. *J. Hosp. Infect.* 56:10–15 (2004).

6. Dancer, S.J.: The role of environmental cleaning in the control of hospital-acquired infection. *J. Hosp. Infect.* 73:378–385 (2009).

7. Boyce, J.M., N.L. Havill, D.G. Dumigan, M. Golebiewski, O. Balogun, and R. Rizvani: Monitoring the effectiveness of hospital cleaning practices by use of an adenosine triphosphate bioluminescence assay. *Infect. Control Hosp. Epidemiol.* 30:678–684 (2009).

8. Carling, P.C., J.L. Briggs, J. Perkins, and D. Highlander: Improved cleaning of patient rooms using a new targeting method. *Clin. Infect. Dis.* 42:385–388 (2006).

9. Lewis, T., C. Griffith, M. Gallo, and M. Weinbren: A modified benchmark for evaluating the cleaning of some hospital environmental surfaces. *J. Hosp. Infect.* 69:156–163 (2008).

10. White, L., S.J. Dancer, C. Robertson, J. McDonald: Are hygiene standards useful in assessing infection risk? *Am. J. Infect. Control* 36:381–384 (2008).

11. Obee, P., C.J. Griffith, R.A. Cooper, and N.E. Bennion: An evaluation of different methods for the recovery of meticillin-resistant Staphylococcus aureus from environmental surfaces. *J. Hosp. Infect.* 65:35–41 (2007).

12. R Development Core Team: R statistical software package, version 2.13.0, (2011). Available: http://www.r-project.org (accessed September 28, 2011).

13. White, L.F., S.J. Dancer, and C. Robertson: A microbiological evaluation of hospital cleaning methods. *Int. J Environ. Health Res.* 17:285–295 (2007).

14. Sherlock, O., N. O’Connell, E. Creamer, and H. Humphreys: Is it really clean? An evaluation of the efficacy of four methods for determining hospital cleanliness. *J. Hosp. Infect.* 72(2):140–146 (2009).

15. Ferreira, A.M., D. de Andrade, M.A. Rigotti, M.V.F. Ferreira: Condition of cleanliness of surfaces close to patients in an intensive care unit. *Rev. Latino-Am.* 19(3):557–564 (2011).

16. Aycicek, H., U. Oguz, K. Karci: Comparison of results of ATP bioluminescence and traditional hygiene swabbing methods for the determination of surface cleanliness at a hospital kitchen. *Int. J. Hyg. Environ. Health.* 209(2):203–206 (2006).

17. Mulvey, D., P. Redding, C. Robertson, et al.: Finding a benchmark for monitoring hospital cleanliness. *J. Hosp. Infect.* 77(1):25–30 (2011).