Modelling the Costs and Effects of Selective and Universal Hospital Admission Screening for Methicillin-Resistant *Staphylococcus aureus*

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Abstract

**Background:** Screening at hospital admission for carriage of methicillin-resistant *Staphylococcus aureus* (MRSA) has been proposed as a strategy to reduce nosocomial infections. The objective of this study was to determine the long-term costs and health benefits of selective and universal screening for MRSA at hospital admission, using both PCR-based and chromogenic media-based tests in various settings.

**Methodology/Principal Findings:** A simulation model of MRSA transmission was used to determine costs and effects over 15 years from a US healthcare perspective. We compared admission screening together with isolation of identified carriers against a baseline policy without screening or isolation. Strategies included selective screening of high risk patients or universal admission screening, with PCR-based or chromogenic media-based tests, in medium (5%) or high nosocomial prevalence (15%) settings. The costs of screening and isolation per averted MRSA infection were lowest using selective chromogenic-based screening in high and medium prevalence settings, at $4,100 and $10,300, respectively. Replacing the chromogenic-based test with a PCR-based test cost $13,000 and $36,200 per additional infection averted, and subsequent extension to universal screening with PCR would cost $311,000 and $232,700 per additional infection averted, in high and medium prevalence settings respectively. Assuming $17,645 benefit per infection averted, the most cost-saving strategies in high and medium prevalence settings were selective screening with PCR and selective screening with chromogenic, respectively.

**Conclusions/Significance:** Admission screening costs $4,100–$21,200 per infection averted, depending on strategy and setting. Including financial benefits from averted infections, screening could well be cost saving.

Introduction

*Staphylococcus aureus* is one of the most common causes of nosocomial and community-acquired infections. Since the 1980s, methicillin-resistant *S. aureus* (MRSA) nosocomial prevalence levels have increased in most countries [1–3]. An estimated 25,100 nosocomial MRSA infections occurred in the US in 2005 [4], and have been associated with higher costs, higher mortality and an increased length of stay than infections with methicillin-susceptible *S. aureus* (MSSA) [5,6].

The low nosocomial prevalence in Scandinavian countries and the Netherlands has been ascribed to stringent policies to control the spread of MRSA. Bootsma et al. have investigated the contribution of different components of the Dutch *Search and Destroy* policy [7], indicating that admission screening can effectively reduce MRSA in high prevalence settings [8]. In clinical studies selective screening on admission to intensive care units (ICUs) or universal screening at hospital admission yielded conflicting results [9–15]. Universal admission screening might be an economically viable option through prevention of MRSA infections and its associated costs [15], but has not been widely adopted because of the presumed high costs associated with testing and subsequent isolation [16].
Several detection tests are now commercially available, each with different test characteristics and costs. The impact and relative importance of a test’s sensitivity, specificity and test delay depend on the screening strategy used and the MRSA prevalence in the catchment population. Here, we used a modeling approach to assist hospital administrators in informed decision making on the implementation of an admission screening strategy.

The objectives were (1) to estimate the costs of screening and isolation per infection averted for various admission screening strategies, (2) to compare two MRSA detection tests within these strategies and (3) to investigate the relative importance of test sensitivity, specificity and test delay. Our analysis focused on the United States.

**Study design**

We performed an analysis of costs and effects of universal and selective MRSA screening at hospital admission, combined with isolation of identified MRSA carriers, over a timeframe of 15 years, using a 3% annual discount rate [17]. We compared strategies both to each other and to a baseline without screening or isolation. The analysis was conducted from a US hospital’s perspective, and costs are reported in US dollars using price levels of the year 2007.

We used a previously published [8] discrete event simulation model developed with C++, reflecting MRSA transmission within hospitals, to estimate the health and economic outcome of screening and isolation. The incremental cost-effectiveness ratio (iCER) of selected strategies was calculated as the difference in screening and isolation costs divided by the difference in infections, of one strategy over another. We also present the average cost effectiveness ratios (aCEERs) for each strategy, calculated as the costs of screening and isolation costs divided by the difference in MRSA infections, relative to a baseline of no screening and no isolation. As our main outcome measure is the investment costs per infection averted, we counted up-front investment costs of screening and isolation (e.g., lab tests and contact precautions), but excluded cost consequences of averting MRSA infection, such as a shorter hospital stay and averted treatment costs. Instead, we compare estimated investment costs with financial benefits of averted MRSA infections.

**Overview of the simulation model**

Below, we present a brief overview of the model, a more detailed account is available elsewhere [8]. Parameter estimates are based on data obtained in the University Medical Center Utrecht, the Netherlands, unless specified otherwise. The model simulates three hospitals, each with 693 beds (36 18-bed wards and 5 9-bed intensive care units (ICUs)] with a 100% bed-occupancy. The mean length of stay was assumed at 3 and 7 days within ICUs and regular wards, respectively (exponentially distributed). Each hospital has a catchment population of 220,000 individuals, of which 20,000 are known ‘high risk’ individuals that have a 10 times higher probability of being admitted to the hospital, compared to the non-‘high risk’ individuals. This leads to an average hospital stay of 50% ‘high risk’ and 50% non-high risk patients. Additionally, ‘high-risk’ patients are characterized by a life expectancy of 20 years versus 78 years for non-high-risk patients. One can think of the high-risk group as elderly together with immunocompromised patients. Unidentified hospitalized carriers have a daily probability of 3% of being detected through conventional microbiological cultures obtained for clinical reasons [8]. Individuals identified as MRSA carrier during a hospitalization are ‘flagged’, so that they are identified as such on the next admission.

MRSA transmission occurs primarily via patient-to-patient transmission mediated by the hands of health care workers (HCWs). The adherence of HCWs to the hand-washing protocol is assumed to be constant over time. Transmission is 20 times more likely to occur within a given hospital unit, compared to transmission between units. Transmission can also occur via HCWs who are colonized in the nose/throat [18]. In a high prevalence setting, this route is set to be 8 times less important as patient-to-patient transmission. Finally, the transmission rate in ICUs is assumed to be 3 times higher (for both routes) compared to other wards, due to more frequent contacts between HCWs and patients and the higher susceptibility of ICU patients. The transmission parameters were calibrated to obtain a steady-state nosocomial prevalence of 15% at baseline (high prevalence).

We used an average daily probability of developing an infection of 0.59% for a hospitalized carrier [19]. Coello et al. report that half of the 68 infections occurred within 12 days. We can derive a daily probability of 0.59% by dividing the number of infection (68/2 = 34) by the total time at risk (479 patients * 12 days = 5748 days). This results in an infection rate of 8.9 per 10,000 bed days at baseline with 15% nosocomial prevalence. Infection status was not explicitly modeled and, therefore, infected patients had the same infectiousness and discharge probabilities as MRSA carriers. We evaluated all screening strategies in a high and medium nosocomial prevalence setting of initially 15% [20–23] and 5% [24], respectively. This prevalence is defined as the percentage of positive findings when performing a cross sectional screening of all patients in the hospital with a perfect test. For the high prevalence setting, the screening program was initiated after a simulation time of 10 years. This period was used to avoid major effects of the exact initial conditions and to reach a steady state nosocomial prevalence of 15%. This prevalence level corresponds to 5.5% prevalence upon hospital admission. In the medium prevalence setting, the simulations were started using a prevalence <1%, and the screening program was initiated when the average nosocomial prevalence in the three hospitals reached 5% for the first time. The outcome of our stochastic model is presented for one hospital with 693 beds, as the mean of 1000 simulations for each strategy over the full timeframe of 15 years. The 2-sided 95% uncertainty intervals (UIs) cover the results observed in 95% of the simulations.

**Baseline**

At baseline there is neither active screening for MRSA nor isolation of identified or suspected carriers. The nosocomial prevalence remained at a steady state of 15% over the entire time frame in high prevalence settings. As a baseline for the medium prevalence setting, we assumed a steady-state prevalence of 5% over the time frame, although without interventions the prevalence would continue to rise to the high prevalence level.

**Admission screening and isolation**

We evaluated ‘Selective’ screening of ‘high risk’ patients and ‘flagged’ patients only, as well as ‘Universal’ screening of all patients. Both strategies were evaluated with a PCR-based test and a chromatographic media-based test (see table 1 for test characteristics). We define test delay as the time between collection of specimens and the reporting of results to the wards, which includes transport and laboratory time. We assumed a test delay of 0.5 day for PCR, and 1.5 and 2.5 days for the chromatographic media-based test after 24 and 48 hours of incubation, respectively. One swab is taken from patients at admission which is subsequently tested for
MRSA, without confirmation by conventional culture techniques. Identified MRSA carriers are isolated in single rooms, and are not decolonized during their hospital stay. We assumed no limits on isolation capacity to allow the peak isolation capacity required for each screening strategy to be determined by the model.

Base-case assumptions

To simulate a regionally implemented MRSA screening policy, all three hospitals in the model are assumed to implement identical screening strategies at the same time. The chromogenic media-based test is evaluated after 24 and 48 hours of incubation. Patients with positive results are isolated at both time points, with the last result after 48 hours being considered final. Pre-emptive isolation, defined as isolation upon readmission for the duration of the test delay until confirmed negative for carriage of MRSA, is limited to ‘flagged’ patients only. Single room isolation is assumed to reduce the risk of transmission by 80% [8].

Scenario analysis

We additionally investigate four alternatives to our base-case assumptions: (1) full pre-emptive isolation, that includes pre-emptive isolation for ‘high risk’ as well as ‘flagged’ patients; (2) the absence of pre-emptive isolation; (3) only 1 out of the 3 hospitals in the model implements screening; (4) screening with a chromogenic media-based test, using only the results after 24 h of incubation.

Cost data

The total investment cost borne by the hospital is assumed to consist of the additional cost of isolation plus the cost of screening. The screening and isolation costs were calculated by multiplying estimated resource use (including labor) by unit prices (table 2) [source: bureau of labor statistics, US department of labor]. The prices of consumables were provided by the manufacturers. The costs of isolation were calculated assuming that facilities for single room isolation are available, thereby excluding the capital costs of building new infrastructure. The isolation costs consist of contact precautions and additional cleaning of the room in case of a positive screening test. The costs of the screening program consist of tests, laboratory labor, laboratory equipment, labor of taking swabs and of a clinical risk assessment when screening selectively (table 2).

Sensitivity analysis

In a one-way sensitivity analysis we investigated the impact of alternately varying the test sensitivity (50–100%), specificity (50–100%) and test delay (0–5 days), on the costs and infections averted. Additionally, we investigated the impact of varying key model parameters on the aCER. The sensitivity analysis was conducted using the strategy selective screening with PCR in a high prevalence setting.

Results

Screening strategies

Relative to baseline, all strategies reduced MRSA prevalence in the first years of screening, yielding prevalence rates below 1% after 15 years (figure 1). The number of patients screened over this period was roughly 200,000 and 400,000 per hospital for selective screening and universal screening, respectively.

Percentages of patients in isolation over time are characterized by a peak at the start of the screening program (figure 1). The peak
percentage of patients in isolation ranged from 6.2% to 9.1% and 2.9% to 5.0% for high and medium prevalence, respectively, and was higher for universal screening than for selective screening (table 3). The annual costs associated with screening and isolation decrease over time, and are shown for ‘Selective PCR’ and ‘Selective Chromogenic’ in a high prevalence setting (figure 2). The screening costs of PCR testing are higher than for chromogenic testing, but these costs are partially offset by the lower costs of isolation of ‘Selective PCR’.

The total number of infections at baseline - over the 15 year timeframe - amounted to 2,753 and 918 for high and medium prevalence, respectively. Of these infections, the number averted by the different screening and isolation strategies ranged from 2,085 to 2,252 and from 622 to 709 for high and medium prevalence, respectively (table 3).

The least costly strategy in terms of the costs per infection averted is ‘Selective Chromogenic’. The investment costs of this strategy in a high prevalence setting are $8.7 m and it averts a total of 2,085 ($2,085/2,753 = 76%) infections compared to baseline (table 3). In a medium prevalence setting, ‘Selective Chromogenic’ costs $6.4 m and averts 622 (622/918 = 68%) infections compared to baseline.

The most effective strategy was ‘Universal PCR’, averting 2,252 (82%) and 709 (77%) infections in high and medium prevalence settings, respectively. This strategy was also the most costly, requiring a total investment of $16.3 m and $15.0 m for high and medium prevalence, respectively.

To visualize comparisons between strategies, we plotted costs and health gains of each strategy (figure 3). In the high prevalence setting, the aCER of selective screening of ‘high risk’ patients with a chromogenic media-based test (‘Selective Chromogenic’), compared to baseline, is $4,100 per infection averted, which is represented by line A. Substituting the chromogenic media-based test by a PCR-based test (‘Selective PCR’), represented by line B, costs an additional $1.6 m and averts 121 more infections, resulting in an iCER of ‘Selective PCR’ compared to ‘Selective Chromogenic’ of $13,000 per additional infection averted. An extension of ‘Selective PCR’ to all patients (‘Universal PCR’), costs an additional $6.1 m and averts an additional 46 infections, resulting in an iCER of $131,000 per infection averted (line C).

In the medium prevalence setting, the aCER - compared to baseline – of screening ‘high risk’ patients with a chromogenic based test (‘Selective Chromogenic’) is $10,300 per infection averted. Substituting the chromogenic media-based test by a PCR-based test (‘Selective PCR’), represented by line B, costs an incremental $2.1 m and averts an incremental 59 infections, resulting in an iCER of ‘Selective PCR’ compared to ‘Selective Chromogenic’, of $36,200 per additional infection averted. The incremental returns on investment strongly diminish with an extension of ‘Selective PCR’ to all patients (‘Universal PCR’), at an iCER of $232,700 per additional infection averted (line C).

Universal screening with a chromogenic media-based test is dominated in both settings by selective screening with PCR (i.e. selective screening with PCR is both cheaper and more effective).
Table 3. Results of screening strategies.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Test</th>
<th>Screening ($m)</th>
<th>Isolation ($m)</th>
<th>Total Investment Cost ($m)</th>
<th>Cases of infection</th>
<th>aCER (Total investment cost $ per infection averted) (95% UI)</th>
<th>Peak isolation capacity required (%)</th>
<th>Patients screened</th>
<th>Time to 50% prevalence reduction (Yrs)</th>
<th>Prevalence after 15 years (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>High</strong></td>
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<tr>
<td>Baseline</td>
<td>None</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2753</td>
<td>0</td>
<td>NA</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Selective</td>
<td>PCR</td>
<td>6.17</td>
<td>4.05</td>
<td>10.22</td>
<td>547</td>
<td>2,206</td>
<td>4,633 (4,477–4,843)</td>
<td>83,774</td>
<td>6.2</td>
<td>200,179</td>
</tr>
<tr>
<td>Selective</td>
<td>Chromogenic</td>
<td>2.87</td>
<td>5.78</td>
<td>8.65</td>
<td>668</td>
<td>2,085</td>
<td>4,149 (3,948–4,442)</td>
<td>119,407</td>
<td>7.2</td>
<td>200,839</td>
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<tr>
<td>Universal</td>
<td>PCR</td>
<td>10.42</td>
<td>5.89</td>
<td>16.30</td>
<td>501</td>
<td>2,252</td>
<td>7,237 (7,000–7,487)</td>
<td>121,681</td>
<td>7.8</td>
<td>375,725</td>
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<tr>
<td>Universal</td>
<td>Chromogenic</td>
<td>4.21</td>
<td>8.15</td>
<td>12.36</td>
<td>622</td>
<td>2,131</td>
<td>5,799 (5,484–6,142)</td>
<td>168,449</td>
<td>9.1</td>
<td>375,739</td>
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<td><strong>Medium</strong></td>
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</tr>
<tr>
<td>Baseline</td>
<td>None</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>918</td>
<td>0</td>
<td>NA</td>
<td>0</td>
<td>0</td>
<td>0</td>
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<tr>
<td>Selective</td>
<td>PCR</td>
<td>5.81</td>
<td>2.71</td>
<td>8.52</td>
<td>237</td>
<td>681</td>
<td>12,508 (11,454–13,677)</td>
<td>55,981</td>
<td>2.9</td>
<td>188,374</td>
</tr>
<tr>
<td>Selective</td>
<td>Chromogenic</td>
<td>2.69</td>
<td>3.69</td>
<td>6.38</td>
<td>296</td>
<td>622</td>
<td>10,257 (9,110–11,819)</td>
<td>76,226</td>
<td>3.3</td>
<td>188,461</td>
</tr>
<tr>
<td>Universal</td>
<td>PCR</td>
<td>10.42</td>
<td>4.61</td>
<td>15.03</td>
<td>209</td>
<td>709</td>
<td>21,195 (19,841–23,347)</td>
<td>95,310</td>
<td>4.3</td>
<td>375,745</td>
</tr>
<tr>
<td>Universal</td>
<td>Chromogenic</td>
<td>4.21</td>
<td>6.18</td>
<td>10.39</td>
<td>271</td>
<td>647</td>
<td>16,056 (14,593–18,106)</td>
<td>127,664</td>
<td>5.0</td>
<td>375,766</td>
</tr>
</tbody>
</table>

1 The number of patient days in isolation.
2 The peak isolation capacity required by the hospital in 97.5% of all simulations.
3 The number of years required to reach a 50% reduction in the nosocomial prevalence.
The cumulative and discounted costs in US$ (2007) and discounted effects for one hospital over 15 years, using base-case assumptions, for a high (15%) as well as a medium (5%) prevalence setting.
NA not applicable; aCER average cost-effectiveness ratio in $ per infection averted, compared to no screening; UI uncertainty interval;
doi:10.1371/journal.pone.0014783.t003
Scenario analysis

Comparing selective screening with PCR using base-case assumptions with the individual scenarios (table 4), shows that extending pre-emptive isolation from ‘flagged’ patients only to all ‘high risk’ patients, averts 35 (+1.4%) additional infections at an additional cost of $3.6 m (+34.9%). The absence of any pre-emptive isolation reduces the number of infections averted by 32 (−1.4%) and costs by $0.4 m (−4.3%). If only one out of the three hospitals implements screening, the total investment costs are $8.8 m (7.8%) higher than the total investment costs of the 3 hospitals in base case scenario, while the number of infections averted in the participating hospital diminishes by 254 (−11.5%) (158 infections are averted in each of the non-participating hospitals). A screening program using only the results of the chromogenic media-based test at 24 h of incubation, reduces the number of infections averted by 211 (−9.6%) and also costs by $3.6 m (−35.7%), compared to PCR-based screening.

Sensitivity analysis

The investment costs and the infections averted of varying test sensitivity and specificity from 50% to 100% with increments of 5%, are shown in figure 4 (left panel). A higher test sensitivity increases the number of infections averted but has very little impact on costs. A higher specificity strongly reduces costs but has a minor impact on health outcome. The slight increase in infections averted with a decreasing specificity is caused by the higher number of patients that are isolated based on a false positive test result and are therefore at lower risk of transmission.

Discussion

Cost savings

The true costs attributable to MRSA infection are unknown and the appropriate method to determine these costs is debated [16]. Reported additional hospital costs of MRSA infection over no infection range from $6,709 to $64,216, depending on the type of infection [6,9,15,25]. Additional hospital costs of MRSA infections over MSSA infections range from $8,327 to $16,738, depending on the type of infection [6,25,26]. Using a recently published estimate of hospital costs ($17,645 translated to US$ 2007) of MRSA infection over no infection [6], we can compare the financial benefits of averted infections to the investment costs per infection.
averted, and estimate the net benefits (figure 6). If the averted hospital costs of infection are real savings to the hospital, all evaluated screening strategies are cost-saving in a high prevalence setting. The net benefit is estimated at $28.7 m for ‘Selective PCR’, and $28.1 m for ‘Selective Chromogenic’, followed by $25.2 m for ‘Universal Chromogenic’ and $23.4 m for ‘Universal PCR’. In a medium prevalence setting, the net benefits are lower; $4.6 m for ‘Selective Chromogenic’ and $3.5 m for ‘Selective PCR’, followed by $1.0 m for ‘Universal Chromogenic’. ‘Universal PCR’ was not cost-saving in this setting with a net benefit of $−2.5 m.

**Additional considerations**

Our scenario analysis confirms that admission screening will be less effective and more costly if neighboring hospitals do not screen...
Table 4. Results of the scenario analysis.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Test</th>
<th>Screening ($m)</th>
<th>Isolation ($m)</th>
<th>Total Investment Cost ($m)</th>
<th>Cases of infection</th>
<th>Cases of infection averted vs. baseline</th>
<th>aCER (Total investment cost $ per infection averted) (95% UI)</th>
<th>Isolation¹</th>
<th>Peak isolation capacity required (%)²</th>
<th>Patients screened</th>
<th>Time to 50% prevalence reduction (Yrs)³</th>
<th>Prevalence after 15 years (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>None</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2753</td>
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<td>NA</td>
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<td>NA</td>
<td>15</td>
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<td>Base-case scenario</td>
<td>PCR</td>
<td>6.17</td>
<td>4.05</td>
<td>10.22</td>
<td>547</td>
<td>2,206</td>
<td>4,633 (4,477–4,843)</td>
<td>83,774</td>
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<td>Scenario 1 Selective –</td>
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<tr>
<td>Full preemptive isolation</td>
<td>PCR</td>
<td>6.17</td>
<td>7.63</td>
<td>13.80</td>
<td>512</td>
<td>2,241</td>
<td>6,158 (5,920–6,406)</td>
<td>157,568</td>
<td>8.3</td>
<td>200,176</td>
<td>3.37</td>
<td>0.22</td>
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<tr>
<td>No preemptive isolation</td>
<td>PCR</td>
<td>6.17</td>
<td>3.62</td>
<td>9.79</td>
<td>579</td>
<td>2,174</td>
<td>4,502 (4,298–4,703)</td>
<td>74,714</td>
<td>5.5</td>
<td>200,178</td>
<td>3.58</td>
<td>0.37</td>
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<td>Scenario 3 Selective –</td>
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<tr>
<td>1 out of 3 hospitals screens</td>
<td>PCR</td>
<td>6.24</td>
<td>4.79</td>
<td>11.02</td>
<td>801</td>
<td>1,952</td>
<td>5,646 (5,232–6,086)</td>
<td>98,900</td>
<td>5.8</td>
<td>202,360</td>
<td>3.50</td>
<td>2.59</td>
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<tr>
<td>Chromogenic media-based test</td>
<td>Chromogenic 24</td>
<td>2.87</td>
<td>3.70</td>
<td>6.58</td>
<td>758</td>
<td>1,995</td>
<td>3,299 (3,076–3,555)</td>
<td>76,543</td>
<td>5.9</td>
<td>200,730</td>
<td>4.23</td>
<td>0.80</td>
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</tbody>
</table>

1 The number of patient days in isolation.
2 The peak percentage of total patients in isolation in 97.5% of all simulations.
3 The number of years required to reach a 50% reduction in the nosocomial prevalence.

The cumulative and discounted costs in US$ (2007) and discounted effects for one hospital over 15 years, for a high (15%) prevalence setting. A CER average cost-effectiveness ratio in $ per infection averted, compared to no screening. UI uncertainty interval.

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more cost-effective than ‘Selective Chromogenic’, and at thresholds of $45 and $106 for high and medium prevalence settings respectively, ‘Selective PCR’ becomes dominant over ‘Selective Chromogenic’ (i.e., less costly and more effective).

Our results contrast with another recent economic analysis [13], which recommended screening with a chromogenic media-based test after 24 hours of incubation over PCR-based screening. An important difference is that this study assumed an equal test delay of 1 day for both tests, and used a sensitivity and specificity for the chromogenic media-based test of 98.0% and 99.8%, respectively, where we have used 76.6% and 98.6%, based upon a recently performed meta-analysis [28].

**Study limitations**

The outcomes from our study depend on the validity of the transmission model. To assess the validity of our model we conducted extensive sensitivity analyses and have provided estimates around our estimated aCER. We did not perform a full probabilistic sensitivity analysis to estimate the impact of the uncertainty in the assumed model parameter values, because the computation time required would be unfeasibly long for the type of model we used. Instead, the impact of varying model parameters was investigated using one-way sensitivity analysis.

Because a model remains a simplification of real life situations, the inherent limitations should be discussed. No limit was set on isolation capacity and it was assumed that all identified carriers were isolated, with corresponding isolation costs. However, this ideal policy will not always be realized [29]. Failure to isolate will reduce the total isolation effectiveness, but will also reduce costs. In our analyses we considered isolation not to be perfect (80% reduction in infectiousness), but costs were always incurred. This will overestimate the costs per infection averted. We assumed an average rate of infection for all carriers, whereas this rate may differ between patients in ICU and in a regular ward.

There are no published estimates on the additional cost (if any) of a patient in a single room versus a semi-private room or a ward [30], and consequently we have omitted these costs from our analysis, as others have done [9,15]. Some authors have included estimates based on construction costs [9], on the maintenance of the additional floor space required [13], or on revenue lost [31]. These approaches can be valid but are strongly determined by local conditions, such as the type of infrastructure, the shared use of isolation facilities for other pathogens and the level of hospital occupancy. Some additional opportunity costs are likely to occur in a hospital operating at near full capacity, due to bed blocking [32]. We have used sensitivity analysis to estimate the impact of additional single room isolation costs.

As our main outcome measure was investment costs per infection averted, our calculations neglect the benefit of patients of not having MRSA. The healthcare utilization costs of treating MRSA infection are driven by the patient’s length of stay. The length of stay varies considerably across hospitals and even between wards in a single hospital. For hospitals with a relatively short length of stay, the screening strategies investigated in this study will result in lower cost savings and lower net benefits than shown in figure 6.

For a more comprehensive determination of cost-effectiveness from the societal perspective, more data is needed on the value of averted infections in terms of the additional survival, quality of life and the costs of MRSA infection, during hospital stay as well as after discharge. One would also hope to include the potential negative effects of isolation on quality of care [33] and possibly the costs of damage to hospital reputations or subsequent litigations.

Yet, with the aforementioned limitations, this analysis provides a
robust estimate of the costs of averting MRSA infection through screening and isolation. Our estimates can be considered in combination with the hospital’s own estimates, e.g. the additional costs of single room isolation and savings of averted infections, to support decision making on cost-effective infection control strategies.

**Conclusions**

Based upon our simulation model, three important conclusions can be drawn related to MRSA admission screening:

1. Excluding any financial benefits from averted infections, the choice of strategy depends on the setting, the costs of isolation and the hospital’s willingness to pay to avert infection. In both settings, selective screening with a chromogenic media-based test is the least costly strategy in terms of the cost per infection averted. More infections can be averted by replacing the chromogenic media-based test with a PCR test, at additional costs. The additional infections that can be averted with universal screening with PCR are relatively costly.

2. The ranking of strategies is sensitive to additional daily costs of single room isolation. At thresholds of $45 and $106, in high and medium prevalence settings respectively, selective screening with PCR becomes dominant over selective chromogenic media-based screening.

3. Assuming $17,645 benefit per infection averted, all evaluated strategies using base-case assumptions are cost-saving with the exception of universal screening with PCR in a medium prevalence setting. The most cost-saving strategies in high and medium prevalence settings are selective screening with PCR and selective screening with a chromogenic media based test, respectively.

**Author Contributions**

Conceived and designed the experiments: GH MB DMB. Analyzed the data: GH MB DG. Wrote the paper: GH MB ML DG DMB. Principal
References


