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**Graphical abstract**

405 nm irradiation

460 nm irradiation

Endogenous porphyrins

ROS quenched by STX

ROS

STX

Bleached STX

O₂

S. aureus
Dual-wavelength photo-killing of methicillin-resistant Staphylococcus aureus

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Introduction

In recent years, the threat of antimicrobial resistance has become one of the most important concerns for public health. Infection outbreaks that result from multidrug-resistant organisms that have emerged remain a significant problem (1, 2). Staphylococcus aureus infections are among the most important causes of skin and soft tissue infections, with methicillin-resistant Staphylococcus aureus (MRSA) being particularly important (3, 4). Therefore, novel and nontraditional approaches must be explored to quell these negative effects. Over the years, antimicrobial blue light (aBL) at a 405-nm wavelength has been emerging as a potential alternative treatment for localized infections (5). The accepted mechanism responsible for the antimicrobial effects of aBL (405 nm) is through excitation of endogenous photosensitizing porphyrins and the subsequent generation of singlet oxygen, resulting in lipid peroxidation, DNA damage, cell wall damage, and cellular apoptosis of microbial cells (5). However, previous studies and preliminary results in our laboratory showed that MRSA is much more tolerant of aBL at 405 nm than most other species (6). Recent findings by other groups have demonstrated the antioxidant properties of staphyloxanthin (STX), which is a membrane-bound carotenoid pigment of MRSA, responsible for its characteristic golden colony phenotype (7, 8). In addition, it has been demonstrated that STX is subject to photolysis through 460-nm light exposure, rendering it more susceptible to H₂O₂-mediated killing (9). Therefore, we theorized that the limited antimicrobial efficacy of aBL (405 nm) we observed with MRSA was a direct result of STX because this is also a known singlet oxygen scavenger. As a result, we hypothesized that treatment of MRSA using 405-nm light could be improved by STX photolysis using preexposure with 460-nm light. Here we report a potentially novel dual-wavelength irradiation approach using the combination of 460-nm and 405-nm light, which exploited the STX photolytic effect of 460-nm light to sensitize MRSA to 405-nm light.
Results

**STX promoted resistance in *S. aureus* to 405-nm light–mediated killing.** In this study, we investigated whether the presence of STX within MRSA is responsible for its inherent resistance to aBL (405 nm). We compared the killing potential of a WT *S. aureus* strain (Pig1) that produces STX and an ΔcrtM isogenic mutant (Pig1ΔcrtM) of the same strain that is deficient in the production of STX. After 108 J/cm² aBL, a 4.14 log₁₀ CFU reduction was achieved in Pig1ΔcrtM, compared with its WT parental strain in which only a 1.09 log₁₀ CFU reduction was observed (P < 0.001; Figure 1). These findings thus demonstrated the ability for STX to promote resistance to aBL at 405 nm.

**Exposure to 460-nm light sensitized MRSA to 405-nm light.** A recent study discovered that 460 nm irradiation can lyse the STX pigment, rendering MRSA more sensitive to ROS-mediated killing (9). Therefore, we sought to determine whether pre-irradiation with 460-nm light was able to attenuate STX-mediated resistance of MRSA to 405-nm light killing. In the strain MRSA USA300, we found that 108 J/cm² exposure of 405-nm light was only capable of inactivating 0.64 log₁₀ CFUs (Figure 2A). However, preexposure to 460-nm light (180 J/cm²) significantly potentiated the killing of MRSA with a 3.58 log₁₀ CFU reduction (P = 0.004; Figure 2A). Irradiation with 460 nm alone, at an equivalent dose, was insufficient to elicit any antimicrobial effects. In a clinical strain of MRSA (AF0003), we also observed a significant improvement in the killing efficacy of 405-nm light with a preexposure of 460-nm light. aBL (405 nm) alone inactivated 1.83 log₁₀ CFUs, whereas the addition of 460-nm light resulted in a 3.23 log₁₀ CFU reduction (P = 0.02; Figure 2B). As with MRSA USA300, 460-nm irradiation alone, at an equivalent dose, was insufficient to elicit any antimicrobial effects on the strain AF0003.

**STX photolysis alone mediated sensitization of MRSA to 405-nm light.** To determine whether the enhancement in MRSA CFU reduction by 460-nm light was solely based on STX photolysis or whether there was another underlying mechanism, Pig1 ΔcrtM and its parental WT strain were exposed to 405-nm light killing. In the strain MRSA USA300, we found that 108 J/cm² exposure of 405-nm light was only capable of inactivating 0.64 log₁₀ CFUs (Figure 2A). However, preexposure to 460-nm light (180 J/cm²) significantly potentiated the killing of MRSA with a 3.58 log₁₀ CFU reduction (P = 0.004; Figure 2A). Irradiation with 460 nm alone, at an equivalent dose, was insufficient to elicit any antimicrobial effects. In a clinical strain of MRSA (AF0003), we also observed a significant improvement in the killing efficacy of 405-nm light with a preexposure of 460-nm light. aBL (405 nm) alone inactivated 1.83 log₁₀ CFUs, whereas the addition of 460-nm light resulted in a 3.23 log₁₀ CFU reduction (P = 0.02; Figure 2B). As with MRSA USA300, 460-nm irradiation alone, at an equivalent dose, was insufficient to elicit any antimicrobial effects on the strain AF0003.

**Enhancement of 405-nm light killing by 460-nm light was dose dependent.** In this study, we determined the effect of increasing radiant exposures of 460-nm light on the effectiveness of 405-nm light. The representative clinical strain of MRSA (AF0003) was used. MRSA was exposed to different radiant exposures of 460-nm light (90 J/cm², 180 J/cm², or 360 J/cm²; reflecting a preexposure duration of 15, 30, and 60 minutes, respectively), before exposing bacteria to 54 J/cm² or 108 J/cm² of 405-nm light. Preexposure to 360 J/cm² of 460-nm light resulted in the most significant killing of MRSA after 54 J/cm² of 405-nm light, with a killing of 2.14 log₁₀ CFUs compared with 405 nm alone, which only reduced the CFUs by 0.36 log₁₀ CFUs (P = 0.002; Figure 4). Conversely, exposing MRSA to 54 J/cm² of 405-nm light after exposure to 90 J/cm² or 180 J/cm² did not result in any significant improvement (P = 0.9). This suggests that the
enhancement of the antimicrobial efficacy of 405-nm light by 460-nm light is dose dependent, which is not surprising because increased photolysis of STX by 360 J/cm² may have increased the sensitivity of MRSA to lower 405-nm doses. When exposure to 405-nm aBL was increased to 108 J/cm², preexposure to 180 J/cm² significantly enhanced killing of MRSA relative to 405-nm light alone (P = 0.02). At a radiant exposure of 90 J/cm² of 460-nm light, however, there was no significant improvement in MRSA killing when the exposure was 108 J/cm² of 405-nm light (P = 0.40). These findings suggest that enhancement of 405-nm light–mediated killing by preexposure to 460-nm light was contingent on the delivered radiant exposure of 460-nm light, in that increasing radiant exposures of 460-nm light resulted in increased susceptibility of MRSA to 405-nm light. We additionally found there to be some increase in intracellular ROS (1.56-fold) when 405-nm light was administered after photolysis by 460-nm light (360 J/cm²) when compared with 405 nm alone; however, this was not found to be statistically significant (P = 0.24; Supplemental Figure 2; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.134343DS1).

Dual-wavelength 460-nm + 405-nm light exposure effectively reduced the viability of MRSA biofilms. In this study, we investigated whether preexposure to 460-nm light improved the efficacy of 405-nm aBL against 48-hour-old MRSA biofilms. The MRSA AF0003 strain was used as the representative strain for all biofilm experiments. The biofilms were initially exposed to 180 J/cm² of 460-nm light immediately before exposing them to 405-nm light at 108 J/cm², 216 J/cm², or 324 J/cm². We found that irradiation of 108 J/cm² of aBL at 405 nm did not result in any antimicrobial effects in the dual-wave-
length irradiation–exposed group, or in the 405-nm light alone–treated group (Figure 5). When the exposure of 405-nm light was increased to 216 J/cm², however, the dual-wavelength irradiation–treated group showed a CFU reduction of 1.85 log₁₀, compared with a 0.73 log₁₀ CFU reduction in the 405-nm light alone–treated group (P < 0.05). When the exposure of 405-nm light reached 324 J/cm², the dual-wavelength irradiation resulted in a 2.72 log₁₀ CFU reduction in MRSA biofilms compared with 405-nm light alone, which inactivated 1.49 log₁₀ CFUs (P = 0.002; Figure 5). Dual-wavelength of 460-nm + 405-nm light significantly improved the treatment efficacy in a mouse skin abrasion infection model. To predict the clinical translatability of the dual-wavelength irradiation approach combining 460-nm and 405-nm light exposure, we investigated its efficacy in a proof-of-principle mouse skin abrasion infection model. Mice were inoculated within an abrasion wound with approximately 10⁷ CFUs of MRSA USA300 and incubated 3 hours before treatment with the dual-wavelength irradiation. When the infected wounds were treated with the dual-wavelength irradiation (460 nm/342 J/cm² and 405 nm/360 J/cm²), a 1.97 log₁₀ CFU reduction was achieved, compared with 405 nm alone at an equivalent radiant exposure, which only reduced the CFUs by 0.28 log₁₀ CFUs (P < 0.0001; Figure 6; n = 7). Treatment with 460-nm light (342 J/cm²) alone did not influence bacterial viability within the wounds. These findings demonstrated the potential of the dual-wavelength (460 nm + 405 nm) irradiation approach as a viable treatment option for wound infections caused by MRSA.

Dual-wavelength exposure of 460-nm + 405-nm light did not result in apoptosis of mouse skin cells. To determine whether the dual-wavelength irradiation by the combination 460-nm and 405-nm light results in cellular apoptosis of mammalian skin cells, the therapeutic dose of the dual-wavelength irradiation used during the in vivo killing studies (460-nm light/342 J/cm² and 405-nm light/t 360 J/cm²) was administered to naive mouse skin tissue before assessing cellular apoptosis using the TUNEL assay. The results showed no presence of apoptotic cells in the treated group immediately after the dual-wavelength irradiation treatment, 24 hours after treatment, or 48 hours after treatment (Figure 7, A–C). The untreated group did not show any evidence of apoptosis (Figure 7D) compared with the positive control (Figure 7E).
Discussion

In this study, we investigated the role of STX in promoting resistance to 405-nm light killing in MRSA. In addition, we explored the STX photolytic effect of 460-nm light exposure as a means of sensitizing MRSA to 405-nm light killing. For the first time to our knowledge, with the use of STX-deficient Pig1ΔcrtM, we confirmed that the presence of STX within MRSA was directly responsible for the relative resistance phenotype to 405-nm light that has been observed in our and other laboratories (6). There have, however, been studies that have illustrated the sensitivity of 405-nm light killing, which conflict somewhat with ours (10). It is possible that variabilities in the relative abundance of endogenous STX may explain these findings. Furthermore, because STX is present in 90% of strains (11), it is feasible that improved efficacy of 405-nm light may be achieved using these STX-deficient strains, although further work is required to corroborate this hypothesis. Given that STX has a carotenoid structure, and is thus a singlet oxygen scavenger, it is unsurprising that its presence would interfere with damage elicited by singlet oxygen (12). Liu et al. fully characterized the role of STX and found it to be a powerful singlet oxygen scavenger (7). They found that the STX-deficient isogenic mutant ΔcrtM was greater than 100-fold more susceptible to singlet oxygen generated through methylene blue–mediated photodynamic therapy (MB-PDT), relative to its parental WT (7). These findings, therefore, strongly support our findings illustrating the role of STX in promoting resistance to 405 nm–mediated killing. In a previous study, it was demonstrated that photolysis using 460-nm light sensitized MRSA to killing by hydrogen peroxide (9), further supporting the role of STX in eliciting resistance to ROS resulting from 405-nm light illumination.

We next sought to explore whether preexposure to 460-nm light would be adequate to overcome the relative resistance observed when MRSA is exposed to 405-nm light. It was demonstrated previously that 460-nm light illumination was capable of STX lysis in MRSA. We found that the degree of STX photolysis achieved by 460-nm light illumination significantly potentiated the killing of MRSA when subsequently exposed to 405 nm, suggesting that the extent of STX lysis was enough to overcome any innate resistance to 405-nm light killing.

Exposure to 460-nm light alone, however, was insufficient to elicit any antimicrobial effects. Our results are consistent with findings from other studies that found 460-nm light did not significantly influence the viability of MRSA in vitro (6,9). In contrast, however, findings from another study demonstrated that at a wavelength of 470 nm, a significant loss of viability may be achieved in S. aureus (13). Furthermore, a recent study found that combining riboflavin with 450-nm light potentiated the killing of MRSA (14). Riboflavin has been shown to possess photosensitizing properties when activated with UV to eliminate microorganisms (15,16). In addition, riboflavin has been found to be effective in combination with 460-nm light for inactivating Listeria monocytogenes (17), which is not surprising given that riboflavin can absorb light at this wavelength (18). Moreover, MRSA can produce riboflavin endogenously depending on the exogenous availability of riboflavin (19). A study by Chaffin et al. found that genes encoding riboflavin biogenesis were not upregulated when riboflavin was adequately available (such as when grown in Luria-Bertani medium; ref. 20); suggesting that
when exogenous riboflavin is limited, MRSA may overproduce riboflavin endogenously. Therefore, it is reasonable to infer that variabilities in the endogenous production of riboflavin may offer an explanation of the conflicting findings achieved by different studies into the antimicrobial effects of 450-nm to 470-nm light.

In addition to antimicrobial killing with the use of light within the blue light spectrum (i.e., 400 nm–470 nm), there have been other light-based modalities that have effectively eliminated organisms. For example, MB-PDT, a commonly used light-based antimicrobial approach (which uses a photosensitizer [MB] that is activated by red light to generate ROS), has been used against an array of bacterial pathogens, inclusive of MRSA in vitro and in vivo (21, 22). A study by Vecchio et al. found that MB-PDT was able reduce the relative luminescence of a bioluminescent variant of MRSA USA300 inoculated onto a burn (30-minute incubation) by approximately 2 log10 CFUs (22). This reduction in MRSA was similar to what we observed after 3 hours of incubation with MRSA USA300 using the 460-nm + 405-nm light approach. The obvious benefit of our 460-nm + 405-nm light approach is that it is strictly light-based and does not require the addition of any exogenous chemical agent. In addition, other light-based strategies using nonvisible light such as UVC (200 nm–280 nm wavelength) have been employed (23). A study by Dai et al. investigated the effectiveness of UVC on mouse abrasion wounds containing S. aureus or Pseudomonas aeruginosa (23). They found that low radiant exposures of UVC (< 3 J/cm²) effectively reduced the bacterial burden of both species (23). Although UVC is an effective approach to eliminate infection, an important consideration is potential side effects on the surrounding mammalian tissue. Because UVC can be absorbed by host DNA, resulting in DNA damage (24), the application of longer wavelengths such as those within the blue light region have been shown to be considerably safer than UVC (24).

We next investigated whether STX photolysis by 460-nm light was predominantly responsible for potentiating the effects of 405-nm light against S. aureus. We found that in a parental WT strain of S. aureus (Pig1), preexposure to 460-nm light significantly improved the antimicrobial effects elicited by 405-nm light. However, in an isogenic ΔcrtM mutant of the same strain (Pig1) that does not produce STX, similar killing efficacies were observed when exposed to 405-nm light irrespective of 460-nm light preexposure. These findings strongly suggest that the improved anti-MRSA efficacy of the dual-wavelength irradiation over 405-nm aBL alone is predominantly due to STX photolysis by 460-nm light, rendering S. aureus more susceptible to 405-nm light killing.

Given that bacterial biofilms are important factors contributing to antimicrobial resistance, as well as the recalcitrant nature of MRSA infections (25, 26), we investigated whether dual-wavelength irradiation combining 460-nm and 405-nm light could be a potential modality for eliminating MRSA biofilms. With
the exposure of 405-nm light above 108 J/cm², we found that dual-wavelength irradiation significantly improved the antimicrobial effects relative to either light wavelength alone. However, no antimicrobial effects were elicited at less than or equal to 108 J/cm² of 405-nm light in either the 405-nm alone group or 405-nm + 460-nm light combination–treated group. A possible explanation could be the presence of the extracellular polymeric substance (EPS) produced during biofilm formation, which may have attenuated light delivery, resulting in higher radiant exposures being required to elicit antimicrobial effects. Alternatively, MRSA cells within the EPS may be less metabolically active, reflecting a persister cell state that renders them more tolerant to antibiotic stress (27), which may explain the higher light dose required to elicit antimicrobial effects; further work is required to substantiate this hypothesis. A study by Becker demonstrated that light at 453 nm can increase the metabolic activity of keratinocytes in vitro, although at very low radiant exposure (28). Therefore, it is possible that 460-nm light may have increased the metabolic activity of persister cells with biofilms, rendering them more susceptible to 405-nm light killing (at 216 J/
109 CFU/mL within PBS before extraction of STX. The suspended bacteria were then centrifuged at 4000 
L loop and suspended in 10 mL PBS. The CFUs were then adjusted to approximately

overnight on BHI agar from 25% glycerol freezer stocks. Subsequently, bacterial colonies were collected of infection. We established that 342 J/cm2 of 460-nm light in combination with 360 J/cm2 of 405-nm light, we investigated its efficacy in a proof-of-principle in vivo

increased metabolic activity is warranted to determine its potential role in persister cell activation.

To predict the clinical therapeutic potential of the dual-wavelength irradiation approach combining 460-
nm and 405-nm light, we investigated its efficacy in a proof-of-principle in vivo mouse skin abrasion model of infection. We established that 342 J/cm2 of 460-nm light in combination with 360 J/cm2 of 405-nm light was enough to inactivate almost 99% (1.97 log10 CFU reduction) of bacteria within the abrasion wounds of mice. Either wavelength of light on its own, at equivalent doses, was insufficient to result in any significant killing of MRSA compared with the untreated control. This study, therefore, demonstrated a proof-of-principle approach illustrating that the combination of 460-nm and 405-nm light may effectively reduce the MRSA burden within mouse wounds. An important limitation of this study is that using the abrasion infection as a model, coupled with the short incubation of the infection, may not necessarily be reflected in a more established or complicated soft tissue infection. Given the low penetrance of 460-nm and 405-nm light through the skin (5) (0.5 mm), we would expect that in the absence of more sophisticated light delivery methods, such as with the use of optical clearing agents (30) or a microneedle array for interstitial delivery (31), the efficacy of combining 460-nm + 405-nm light may be limited.

We additionally found no evidence of apoptosis (immediately after treatment, 24 hours later, or 48 hours later) resulting from combining 460-nm + 405-nm light against naive mouse skin at a dose required to inactivate approximately 99% of MRSA within a mouse skin abrasion wound, which suggests the therapy may be safely administered. It is important to appreciate, however, that the doses required to significantly inactivate MRSA within more established wound infections may necessitate higher therapeutic doses. In addition, the results only present a qualitative assessment of safety denoted by the presence (or absence) of apoptotic cells, thus suggesting that further studies are warranted to quantitatively evaluate the safety of using 460-nm + 405-nm light against the skin. In conclusion, 460-nm + 405-nm light combination therapy may offer an effective and safe approach for the treatment of MRSA wound infections.

Methods

Blue light sources. Irradiations of 460-nm and 405-nm light were delivered using 2 light-emitting diodes (LEDs; M405L2 and M470L1; Thorlabs) with peak emissions of 460 nm and 405 nm, respectively, and a full width at half-maximum of 25 nm. The irradiance was regulated by altering the distance of the light source aperture and the target with the use of a PM100D power/energy meter (Thorlabs).

Bacterial strains and growth conditions. The bacterial strains used in this study were clinical strains of an MRSA AF0003 strain that was isolated from an infected soldier deployed in Afghanistan and an IQ00064 strain that was isolated from an infected soldier deployed in Iraq, in addition to a bio luminescent MRSA USA300 strain (23) and an MRSA Pig1 strain (7). The isogenic ΔcrtM mutant from the S. aureus Pig1 strain (7) was used for mechanistic studies. With the exception of the S. aureus Pig1 ΔcrtM strain, all are STX producers (Supplemental Figure 1). The bacteria were cultured in brain-heart infusion (BHI) medium (agar or broth) at 37°C or in an orbital incubator (37°C; 180 rpm).

Extraction of STX from S. aureus strains. All strains of S. aureus used in the study were initially grown overnight on BHI agar from 25% glycerol freezer stocks. Subsequently, bacterial colonies were collected with the use of a 10 μL loop and suspended in 10 mL PBS. The CFUs were then adjusted to approximately 106 CFU/mL within PBS before extraction of STX. The suspended bacteria were then centrifuged at 4000 × g for 4 minutes to pellet the cells. The supernatant was then discarded, and the pellet was suspended in 2 mL of 100% methanol and incubated at 55°C for 3 hours to ensure complete extraction of the pigment. After incubation, the cells were pelleted and the supernatant containing STX was transferred to a 1 mL cuvette. The absorption was then measured using an Evolution 300 UV-Vis Spectrophotometer (Thermo Fisher Scientific) in accordance with the manufacturer’s instructions.

Light killing (460 nm + 405 nm) of planktonic MRSA in vitro. Bacteria were cultured to the mid-log phase and adjusted to approximately 108 CFU/mL in PBS and transferred to a 35 × 12 mm dish. When necessary, bacteria were pre-irradiated with 460-nm light (90–360 J/cm2/100 mW/cm2), before 405-nm light (108 J/cm2/60 mW/cm2) exposure. During light irradiation, the bacterial suspension was stirred using a 12-mm magnetic bar (20 rpm) to ensure uniform exposure of cells to aBL. Aliquots (40 μL) of the suspension were
then withdrawn at varying time points after the initiation of 405-nm light, and the CFUs were determined by serial dilution (10⁻²–10⁻⁷ dilution factors) on BHI agar plates as described previously (32). Experiments were performed in triplicate (3 independent replicates spanning 3 days).

**Measuring intracellular ROS after 460-mm + 405-mm light exposure.** Bacteria were cultured to the mid-log phase and adjusted to approximately 10⁶ CFU/mL in PBS, and 200 μL were transferred to a 96-well plate. The suspension was irradiated with 360 μJ/cm² of 460-mm light, and then 10 μM of CellROX Green Reagent (Thermo Fisher Scientific) was added to the bacterial suspension and incubated for 30 minutes to ensure permeabilization of the probe into cells. The bacterial suspension containing the probe was subsequently irradiated with radiant exposures of 3.6 J/cm², 7.2 J/cm², or 14.4 J/cm² of 405-mm light (low radiant exposures selected due to limited photostability of the probe) before measuring relative fluorescence emission using a SpectraMax M5 multimode multiplate reader. The excitation/emission settings were set to 485/520 nm. Untreated bacteria, 405 nm, and measurements following 460-mm treatment (reflecting the time of 405-mm exposure) were run in parallel. Experiments were performed in triplicate (3 independent replicates spanning 3 days).

**Light killing (460 mm + 405 nm) of MRSA biofilms.** Bacterial suspensions in BHI broth were incubated in 96-well microtiter plates (200 μL/well; approximately 10⁶ CFU/mL), as described previously, for 48 hours to induce biofilm formation (26, 27). Following incubation, the wells were rinsed 3 times with 200 μL PBS to ensure removal of residual medium or planktonic cells, while ensuring that the adhered biofilms remained intact. Before light irradiation, aliquots of 200 μL fresh PBS were added to the wells. The biofilms were pre-irradiated with 180 J/cm² of 460-mm light at an irradiance of 100 mW/cm² immediately before 405-mm light exposure (60 mW/cm²) until radiant exposures of up to 324 J/cm² were delivered. The adhered cells within the biofilms were then isolated by thorough scraping the appropriate wells with a pipette tip and transferring the 200 μL cell/biofilm suspension to a 1.5 mL microcentrifuge tube. A further 200 μL of PBS was added to the wells, and the biofilm isolation was repeated in triplicate to ensure adequate removal of bacterial biofilms. The collected cells were subsequently sonicated using a Branson 2510 Ultrasonic Cleaner (Marshall Scientific, LLC) to ensure that cells within biofilms became adequately dislodged for subsequent plating and CFU quantification. Experiments were conducted in biological triplicate (3 independent replicates spanning 3 days).

**Inoculation of MRSA within a mouse skin abrasion wound.** Mice (Female BALB/c) aged 6–8 weeks and weighing approximately 17–19 g were procured from Charles River Laboratories. Before producing the skin abrasion wounds in mice, mice were injected intraperitoneally with the use of a ketamine/xylazine cocktail (20 mg/kg–100 mg/kg). The mice were then shaved, and the tissue was carefully abraded within a defined 1.0-cm × 1.0-cm area using a no. 15 sterile scalpel blade. The scraped area either did not produce any blood or bled very little. Immediately after the abrasion, 100 μL of PBS were added to the wells. The biofilms were pre-irradiated with 180 J/cm² of 460-mm light at an irradiance of 100 mW/cm² immediately before 405-mm light exposure (60 mW/cm²) until radiant exposures of up to 324 J/cm² were delivered. The adhered cells within the biofilms were then isolated by thoroughly scraping the appropriate wells with a pipette tip and transferring the 200 μL cell/biofilm suspension to a 1.5 mL microcentrifuge tube. A further 200 μL of PBS was added to the wells, and the biofilm isolation was repeated in triplicate to ensure adequate removal of bacterial biofilms. The collected cells were subsequently sonicated using a Branson 2510 Ultrasonic Cleaner (Marshall Scientific, LLC) to ensure that cells within biofilms became adequately dislodged for subsequent plating and CFU quantification. Experiments were conducted in biological triplicate (3 independent replicates spanning 3 days).

**Light killing (460 nm + 405 nm) of MRSA within mouse skin abrasion wounds.** Three hours after inoculation of mouse abrasion wounds, MRSA was treated within mouse wounds. During light exposure in vivo, an adjustable collimator was attached to each light source, 460-nm and 405-nm light, and the LEDs were held 16 cm from the target, angled at an approximately 45° angle, and the collimated light beam was adjusted to cover the abrasion wound (see Figure 8). The 460-nm light was initially delivered onto the infected wound for 15 minutes (200 mW/cm²) before irradiation with 405-nm light (60 mW/cm²). After the 405-nm light was initiated, the 460-nm light irradiance was lowered to 30 mW/cm² and exposed continuously during 405-nm light illumination (90-minute duration). The purpose of continuous exposure of 460-nm light was to ensure that continuous photolysis occurred during 405-nm illumination, as it was found that in a medium that supports MRSA growth (i.e., in vivo), STX can become replenished when 405-nm light exposure ceases (9). The total radiant exposure of 460-nm light was 342 J/cm² and 405-nm light was 360 J/cm² (reflecting a 105-minute treatment time). A total of 7 mice were included for each group, and experiments were performed over 7 days (1 mouse for each group per day).

**Quantification of bacterial CFUs in mouse wounds after light treatment.** Immediately after treatment, mice were euthanized in accordance with standard procedures, and the 1-cm × 1-cm infected skin tissue was isolated and homogenized in 1 mL PBS within a FastPrep lysing matrix tube A (MP Biomedicals) in a FastPrep-24 Classic Instrument (MP Biomedicals). Samples were homogenized for 5 minutes, with 60 sec-
ond intervals, and samples were placed on ice for 5 minutes in between homogenization cycles to limit heat generation. After homogenization, the CFUs were determined by serial dilution (10^3–10^7 dilution factors) on BHI agar plates as described previously (32).

**TUNEL assay to detect apoptotic cells in mouse skin treated with 460-nm + 405-nm light.** The presence of apoptotic cells that resulted from the dual-wavelength irradiation therapy was determined in healthy mouse skin as described previously (33). In brief, skin from the mouse was isolated at 0 and 24 hours, following the treatment with 460-nm light (342 J/cm²) and 405 nm (360 J/cm²). An untreated skin sample was also included as the control and was immediately fixed in 10% phosphate-buffered formalin (Thermo Fisher Scientific) for 48 hours. Tissue sections (4 μm thick) were then analyzed with the DeadEnd Fluorometric TUNEL System (Promega) in accordance with the manufacturer's instructions. Fluorescence images were visualized with the use of the NanoZoomer S60 digital slide scanner where a fluorescein isothiocyanate was used as the fluor and DAPI as the nuclear counterstain. In addition, a section treated with DNase I (RQ1 RNAsefree DNase, Promega), which induces significant DNA damage (34), served as a positive control mouse skin. Fluorescence images were visualized with the use of the NanoZoomer S60 digital slide scanner where a fluorescein isothiocyanate was used as the fluor and DAPI as the nuclear counterstain. In addition, a section treated with DNase I (RQ1 RNAsefree DNase, Promega), which induces significant DNA damage, served as a positive control.

**Statistics.** Data were presented as the mean ± SEM, with differences between means being compared for significance, where appropriate, by either a paired t test or 1-way ANOVA. P values of less than 0.05 were considered significant.

**Study approval.** All animal procedures were approved by the Institutional Animal Care and Use Committees of Massachusetts General Hospital (protocol number: 2015N000187) in accordance with NIH guidelines.

**Author contributions**

TD proposed the concept of dual-wavelength photo-killing of MRSA. LGL and XG established a proof-of-principle for the therapeutic modality. LGL performed in vitro and in vivo studies. JXC and DCH helped provide mechanistic insights into the dual-wavelength photo-killing approach, aiding in the study design. TD and LGL designed experiments and wrote the paper.

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