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## Bacteria killing in ICU associated infections: antibacterial nanosheets as disinfectant†

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Here we report hybrid antibacterial nanosheets (Zn–CuO@GO) where graphene oxide (GO) is decorated with zinc-doped copper dioxide (Zn–CuO) nanoparticles and applied as disinfectant agents to combat multidrug resistant bacterial strains from intensive care units (ICUs). The difference between different Zn–CuO decoration ratios was carefully evaluated and the results indicated when Zn–CuO was deposited onto GO at 4 : 1 weight ratio (w/w), the obtained Zn–CuO@GO exhibited the best dispersity, which was suitable as an antibacterial supplement for disinfectant. Importantly, the Zn–CuO@GO (4 : 1, w/w) based disinfectant could actively kill 100.0% multidrug bacteria strains within 10 min, including multidrug resistant (MDR) *E. coli* and methicillin resistant *S. aureus* (MRSA). Moreover, we measured the antibacterial activities of Zn–CuO@GO at other decoration ratios (i.e., 2 : 1 and 1 : 1, w/w) to study the effect of Zn–CuO decoration density on the antibiosis activity of Zn–CuO@GO. It was also found that decreased Zn–CuO deposition density not only caused the agglomeration of Zn–CuO@GO nanosheets (largely due to the relatively increased GO ratio), but also extended their required function time to 30 min for complete bacterial killing. Taking Zn–CuO@GO (4 : 1, w/w) as the optimized antibacterial nanosheet, further TEM, SEM and LCSM images revealed that Zn–CuO@GO functions via penetrating and wrapping into the cell wall, inducing bacterial sedimentation and cytoplasma leakage, which make them promising for next-generation disinfectant substitution.

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## Introduction

Disinfectant agents are essential components in the practice of control and prevention of bacterial infections, especially in intensive care units (ICUs), where there have been extensive reports on multidrug resistant bacteria such as carbapenem-resistant *Acinetobacter baumannii*, methicillin resistant *S. aureus* (MRSA) and multidrug resistant (MDR) *E. coli*, etc.<sup>1a</sup> Traditionally, iodophor and hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) were widely used as ICU disinfectant due to their strong oxidizing abilities to cause the denaturing of bacteria and consequently bacterial progressive lysis.<sup>1b,c</sup> Given the fact that ICU associated bacteria could develop mutations to convert to multi-drug resistance, Kempf *et al.*, demonstrated that iodophor and H<sub>2</sub>O<sub>2</sub> resistant microorganism emerged recently, and these multi-drug resistant bacteria were growing more invulnerable.<sup>1d</sup>

Additionally, Campos *et al.*, have recently found that bacteria (e.g., MRSA) was resistant against a variety of other candidate disinfectants including sodium hypochlorite, glutaraldehyde, formaldehyde, ethanol, chlorhexidine gluconate, and quaternary ammonium.<sup>1e</sup> A practical method to combat against bacterial resistance is thus urgently to be developed.

The application of nanomaterials into bacterial killing is a promising alternative solution, since these nanoparticles have been proven to be highly reactive in antibiosis compared with those chemical compounds.<sup>2a,b</sup> Several mechanisms, including reactive oxygen species (ROS), metal ion leaking, electrostatic interactions and structural associated physical damage,<sup>2c,d</sup> are proposed to contribute to their antibacterial behaviours. Among them, physical damage to bacteria is irreversible, and this class of materials emerges as ideal candidates for the development of disinfectants. One of the promising nanomaterials is graphene oxide (GO), a strongly oxygenated and highly hydrophilic two dimensional (2D) layered material with biocompatibility, large specific surface area, high chemical and thermal stability,<sup>3a-c</sup> and most importantly, bacterial killing abilities. Fan *et al.*, have recently introduced a GO based antibacterial paper which exhibited minimal cytotoxicity to mammalian cells, but effective towards bacterial inhibition.<sup>3d</sup> Elimelech *et al.*, further demonstrated that GO might cause bacterial lysis by means of their sharpened edges and elevated membrane pressure.<sup>3e</sup> Similarly, Ivanova *et al.*, proposed that GO inactivated

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microorganism through the density and orientation of their graphene edges, which led to pore forming in bacterial membrane.<sup>3f</sup> Despite these recent achievements, GO itself, however, was rarely evaluated against multi-drug resistant bacteria, such as multidrug resistant (MDR) *E. coli*, methicillin resistant *S. aureus* (MRSA).<sup>4a-c</sup> Moreover, GO were typically evaluated on assembled GO integrity, including GO paper<sup>3e</sup> and GO membrane,<sup>3f</sup> while antibacterial effect from individually dispersed GO nanosheet was unsatisfying.<sup>3c</sup> Strategy to decorate GO nanosheets with structure-featured metal oxides that share the physical damage towards bacteria was recently proposed to address this issue, as it not only took advantage of the large specific surface area and morphological features from GO, but also introduced the bacterial activities of metal oxides simultaneously.<sup>4d-f,5</sup>

Previously, Wu *et al.*, have proposed a Zn–CuO@GO decorated commercial porous nickel (Ni) electrode and demonstrated its ability for efficient capture, rapid killing and ultrasensitive detection of bacteria (normal bacterial stains such as *E. coli* (ATCC 25922) and *S. aureus* (ATCC 29213) was evaluated).<sup>6a</sup> Its behaviour upon contact with multi-drug resistant strains remained elusive, and despite of its effectiveness, the electrode was not practical to apply into ICU-associated disinfectant due to its size (6 mm × 24 mm). In contrast, we were able to synthesis Zn–CuO@GO nanosheets to apply as disinfectants and demonstrated their activities to combat against multi-drug resistant bacteria strains. Different from previous works, we obtained Zn–CuO@GO nanosheet solely through *in situ* deposition and growth, and also, we prepared Zn–CuO@GO with different deposition ratios (*i.e.*, Zn–CuO : GO = 4 : 1, 2 : 1 and 1 : 1, w/w) where Zn–CuO that possessed a prickly architecture was allowed to deposit and grow *in situ* onto GO nanosheets to obtain the hybrid antibacterial nanosheets.<sup>6b,c</sup> The Zn–CuO@GO suspensions were applied as a disinfectant. Taking multi-drug resistant *E. coli* (MDR *E. coli*) and a methicillin resistant *S. aureus* (MRSA) as examples, these antibacterial nanosheets could effectively inhibit bacterial growth *via* physical damage, which was confirmed through TEM and SEM. Results indicated that these hybrid nanosheets function as effective antibacterial agents to penetrate into the cellular wall and cause bacterial lysis upon contacting, which was largely due to the fact that they possessed the ‘blades’ from GO and ‘pierces’ from prickly Zn–CuO simultaneously.<sup>7a</sup> As these antibacterial nanosheets function to cause bacterial lysis through physical damage, possible genetic mutation and development of other drug resistant mechanism might not be applicable.<sup>7</sup> To this end, antibacterial Zn–CuO@GO nanosheets is promising as an alternative disinfectant applied in ICU.

## Results and discussion

By taking advantage of sonochemical method, we fabricated GO nanosheets packaged with prickly Zn–CuO. Briefly, the metal ions ( $\text{Cu}^{2+}$  and  $\text{Zn}^{2+}$ ) were firstly allowed to chelate onto GO nanosheets, where ultrasmall nanoclusters were then deposited and readily grew *in situ* *via* sonochemical assisted green energies (Fig. 1A).<sup>7a</sup> As indicated in the transmission electron microscopy

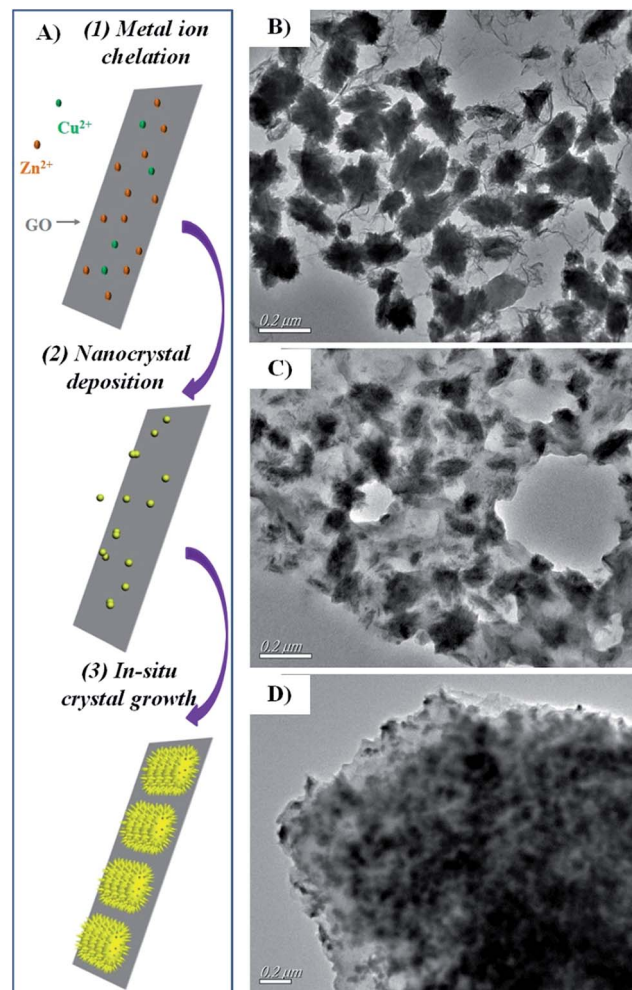


Fig. 1 (A) Schematic illustration of the formation of Zn–CuO@GO from sonochemical method. TEM image of Zn–CuO@GO (4 : 1) (B), Zn–CuO@GO (2 : 1) (C), and Zn–CuO@GO (1 : 1) (D).

image (TEM, Fig. 1B), the prickly Zn–CuO were randomly and successfully deposited onto individual GO nanosheets (4 : 1, w/w), which was similar to previous reports (specifically, the deposition weight ratio was calculated from ICP results (Table S1, ESI†)). The dark rugby-like shape and the sublayer sheet indicated for the prickly Zn–CuO and GO, respectively. As indicated in Fig. 1B, the pricks from Zn–CuO were well preserved, arising from their successful *in situ* nanocrystal deposition and nanoparticle growth. Different from previous works, we obtained Zn–CuO@GO solely from the sonochemical reactions other than scratched from the electrode, and prepared hybrid antibacterial nanosheet in other deposition ratios, *i.e.*, 2 : 1 (Fig. 1C) and 1 : 1 (Fig. 1D) (w/w) for comparison. Compared with Zn–CuO@GO obtained at 4 : 1 (w/w), Zn–CuO@GO prepared at 2 : 1 and 1 : 1 (w/w) were largely in agglomerate (Fig. 1C and D). It is possibly due to the comparably overabundance of GO nanosheets, as there exists strong  $\pi$ – $\pi$  stacking between GO nanosheets. Also, the zeta potential of the hybrid antibacterial nanosheet was measured in saline (ESI, Fig. S1†), which indicated that it was positively charged after Zn–CuO ( $\zeta > 0$ ) was deposited into GO nanosheets ( $\zeta < 0$ ).



As indicated in Fig. 2, compared with Zn–CuO (Fig. 2a) and GO (Fig. 2b), the typical XRD peaks of the hybrid antibacterial nanosheet (Fig. 2c–e) exhibited similar pattern with pristine prickly Zn–CuO at  $2\theta = 32.47^\circ$ ,  $35.49^\circ$ ,  $38.68^\circ$ ,  $48.65^\circ$ ,  $58.25^\circ$ , and  $61.45^\circ$  (Fig. 2a–c), indicating the existence of Zn–CuO in the GO layers.<sup>7a</sup> The reason why there were no obvious peaks from GO that may bear a typical peak at  $12^\circ$  (Fig. 2b) was possibly due to the coverage of other strong peaks from Zn–CuO NPs, which was also in consistence with previous reports.<sup>4g</sup> In synergistic with TEM images in Fig. 1B–D, the synthesis of Zn–CuO@GO was demonstrated successful.

Further, the biocidal activity of Zn–CuO@GO suspension based disinfectant was quantitatively evaluated against a multi-drug resistant *E. coli* (MDR *E. coli*) and a methicillin resistant *S. aureus* (MRSA).<sup>7a</sup> Briefly, after treatment with Zn–CuO@GO suspensions (at concentrations of  $0.1 \text{ mg mL}^{-1}$ ), the viable bacteria were monitored by counting the number of colony-forming units (CFU) with respect to time.<sup>7a</sup> As presented in Fig. 3, representative results clearly indicated that Zn–CuO@GO effectively inactivated those multi-drug resistant strains nearly completely after 10 min of treatment (yellow and green dashed lines), except for Zn–CuO@GO (1 : 1) (purple dashed lines). Compared with GO which exerted minimal cytotoxicity towards these bacterial strains even after over 3 h of treatment (blue bar), it could be concluded that after integrated with GO sheets, the deposited prickly Zn–CuO could greatly enhance their combinational bacterial killing activity.<sup>4f,7a</sup> Notably, the anti-bacterial abilities of Zn–CuO@CuO at different mass ratios were different, as Zn–CuO@GO (4 : 1) and Zn–CuO@GO (2 : 1) shared similarity in superiority in combating multi-drug resistant microbes, while Zn–CuO@GO (1 : 1) took more time ( $\sim 30 \text{ min}$ ) for a complete bacterial inhibition. Based on a 10 min time scale after treatment, GO might account for a maximum 3.6% antibacterial activity while Zn–CuO accounted for at least 96.4%. Thus, we concluded that the deposition density of Zn–CuO contributed significantly to the combinational antibacterial activities of Zn–CuO@GO, and Zn–CuO@GO (4 : 1) exhibited the best performance. Additionally, taking consideration of its flexible architecture and good

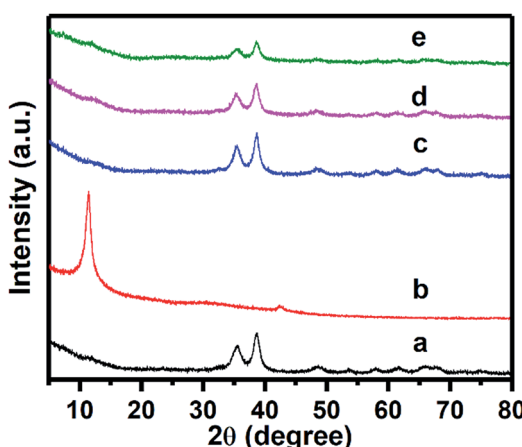


Fig. 2 XRD pattern of Zn–CuO (a), GO (b), Zn–CuO@GO (4 : 1) (c), Zn–CuO@GO (2 : 1) (d), and Zn–CuO@GO (1 : 1) (e), respectively.

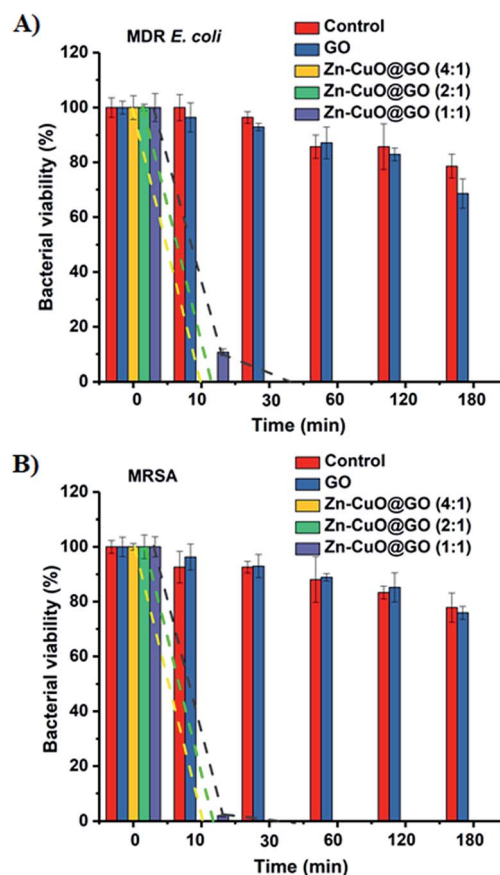


Fig. 3 Bacterial viability after treatment of GO, Zn–CuO@GO (4 : 1), Zn–CuO@GO (2 : 1), and Zn–CuO@GO (1 : 1) at  $0.1 \text{ mg mL}^{-1}$  for different time intervals, respectively. Results for MDR *E. coli* and MRSA were represented in (A and B), respectively.

dispersity, which introduced many possibilities for further biomedical application,<sup>7b,c</sup> the Zn–CuO@GO (4 : 1) was recognized and chosen as the optimized composite for further evaluation. If not otherwise mentioned, the following Zn–CuO@GO refers to Zn–CuO@GO (4 : 1). In comparison with selected works on similar materials (Table S2, ESI†), it was suggested Zn–CuO@GO actively inhibited multi-drug resistant bacteria stains almost completely within 10 min, which rendered themselves promising candidates as a ICU applicable disinfectant.

Live/dead dual-staining was conducted to investigate the bacterial viability after exposure to Zn–CuO@GO, and bacterial without any treatment was chosen as control (Fig. 4).<sup>4a,7c</sup> PI was used specifically to stain membrane perturbed cell with red fluorescence and DAPI was used to label all bacterial nucleus with blue fluorescence. Zn–CuO@GO was also pre-stained with FITC, a green fluorescence utilized widely, to indicate its existence. Notably, bacteria were stained with PI before fixation to indicate the ratio of dead bacteria. It was found that there was scarcely red fluorescence detected for both MDR *E. coli* and MRSA without treatment, which was in consistence with previous reports. After treated with Zn–CuO@GO at  $0.1 \text{ mg mL}^{-1}$ , however, it was found that there were severe aggregations



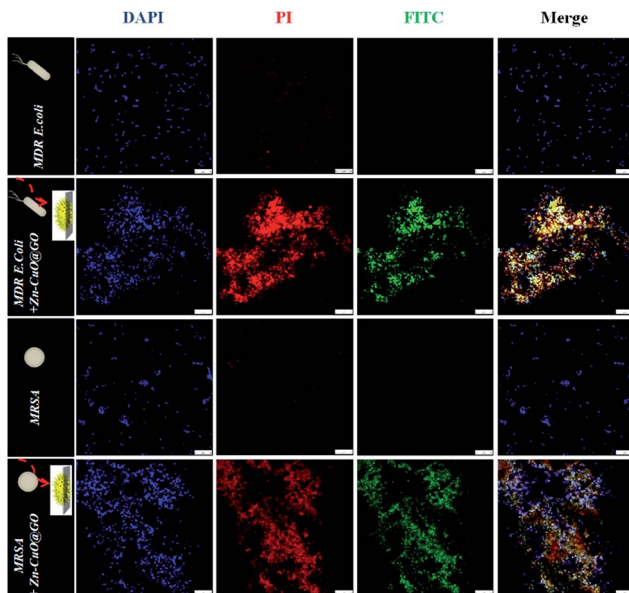


Fig. 4 The laser confocal scanning microscope images of MDR *E. coli* and MRSA before and after 30 min treatment with Zn–CuO@GO nanosheets. The concentrations of Zn–CuO@GO were fixed at  $0.1 \text{ mg mL}^{-1}$  in 0.9% NaCl aqueous solution. DAPI (blue fluorescence), PI (red fluorescence), and FITC (green fluorescence) were used to stain total bacteria, dead bacteria, and the Zn–CuO@GO composite, respectively. Scale bar is  $10 \mu\text{m}$ .

observed, and almost all bacteria were dead as they were stained with strong red fluorescent. The merged images where blue, red and green fluorescent almost fitted together further suggested that compared with previous reports,<sup>7a</sup> the treated bacteria were largely accumulated and adsorbed into the Zn–CuO@GO, where their green fluorescent was in good coincidence with bacteria. This phenomenon demonstrated that after integrated with GO sheets, Zn–CuO could effectively cause the accumulation of bacteria onto their recombinant sheet structure to accelerate bacterial lysis.

To elucidate the bacteria–material interactions after treatment, we first tested whether possible antibacterial mechanisms from zeta potential, metal ions and ROS was dominant. Assuming the fact that if zeta potential was dominant in contributing to the antibacterial activities, Zn–CuO@GO ( $\zeta > 0$ , Fig. S1, ESI†) should have better activity against Gram-negative MDR *E. coli*, we carefully examined the antibacterial results in Fig. 3 again. The only clue we could detect, however, was that Zn–CuO@GO might be more effective against Gram-positive MRSA (Fig. 3), given that Zn–CuO@GO (1 : 1) possessed a better activity towards this strain after 10 min treatment. Such a difference was not detectable for Zn–CuO@GO (4 : 1 or 2 : 1), as they could give a complete inhibition of both strains within 10 min and exhibited no difference. It was thus indicated that zeta potential might not be a determining factor. Further, we examined the effect from trace amount of metal ion leakage towards bacterial inhibition. As provided in Table S3 (ESI†), however, the time-lapse release of metal ions in Zn–CuO@GO indicated that, there were minimal leakage of metal ions such as  $\text{Cu}^{2+}$  and  $\text{Zn}^{2+}$  (only  $1.2 \text{ mmol L}^{-1}$  of  $\text{Cu}^{2+}$  and

$0.004 \text{ mmol L}^{-1} \text{ Zn}^{2+}$  was detected over 3 h of treatment). The trace amount of  $\text{Cu}^{2+}$  and  $\text{Zn}^{2+}$  into surrounding environment in turn suggested the leakage of metal ions was hardly a determining factor for Zn–CuO@GO associated bacterial killing. Additionally, the effects of oxidative stress and ROS production were also tested. For ROS production, a ESR spinning technique was applied, which was effective towards the detection of ROS signals. An important part in ROS generation, OH radicals ( $\cdot\text{OH}$ ), however, was not detectable here under the actual doses of  $0.1 \text{ mg mL}^{-1}$  even upon contact with bacteria. From the ESR curves in Fig. S2,† the arrow pointed area, which should be indication of the existence of ROS, gave no significant difference even after treatment. To advance our knowledge of possible oxidative stress introduced by Zn–CuO@GO, we also took an *in vitro* GSH (glutathione) oxidation assay, of which GSH was an antioxidant in bacteria that could prevent damages to cellular components caused by oxidative stress.<sup>7d</sup> Through examination of loss of GSH concentration after incubation with Zn–CuO@GO (ESI, Fig. S2†), we found that after 2 h of treatment there were no evident difference in the GSH loss. In fact, loss of GSH was minimal (Zn–CuO@GO (~23.45%) and GO (~21.35%)) in comparison with  $\text{H}_2\text{O}_2$  (100%), indicating that Zn–CuO@GO was not functioning through oxidative stress mechanism. Taken together, we have concrete reasons to believe that Zn–CuO@GO cause severe bacterial lysis differently.

Given its unique nanostructure, we then evaluated physical damage associated antibacterial mechanism, and scanning emission microscope (SEM) was first applied to give detailed elaboration on morphological changes of bacterial after co-incubating with Zn–CuO@GO in saline (Fig. 5). It was found that bacteria without treatment maintained their intact membrane without significant morphological changes (ESI, Fig. S3†). After 30 min extensive contact with Zn–CuO@GO, however, it was observed that Zn–CuO@GO nanosheets were attached tightly onto bacterial membrane, which was similar to Zn–CuO alone. It was shown in Fig. 5A and B, that the bacteria lost their cell membrane integrity, as indicated by their severe

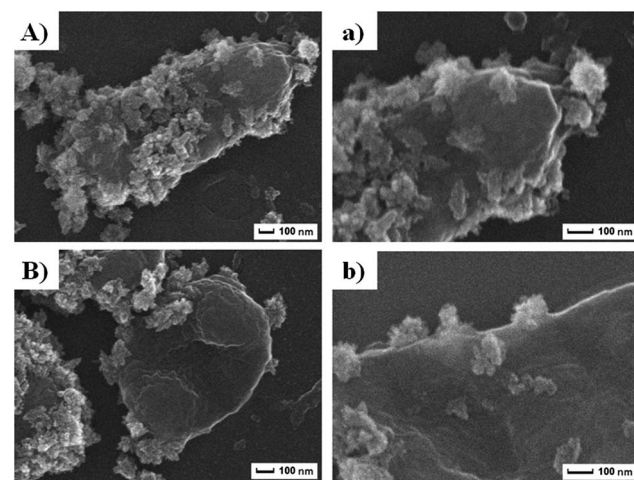


Fig. 5 (A and B) were SEM images of MDR *E. coli* and MRSA after treatment with Zn–CuO@GO, respectively. (a and b) were magnified SEM images revealing detailed cell–material interactions.



shrinkage, which was especially obvious in Fig. 5B. Notably, Zn–CuO@GO may act similarly but in a more effective way taking advantage of Zn–CuO and GO together. Through closer examination of the bacterial surface after treatment, it was found that the surface of Zn–CuO treated bacteria was largely corrugated due to their lost of membrane integrity (ESI, Fig. S4†), while that from Zn–CuO@GO treated bacteria was smoother (Fig. 5a and b), which was attributed to the attachment of Zn–CuO@GO – a layered structure that covered the bacterial membrane. As illustrated in Scheme 1(I and II), in linear with prickly Zn–CuO NPs alone (I), the deposited Zn–CuO NPs on Zn–CuO@GO nanosheets enable them to effectively anchor into the cell membrane (II), and their delicate nano-pricks could then penetrate into the cell walls from diverse positions, causing the leakage of cytoplasm and subsequent bacteria lysis, which was evidenced by the existence of Zn–CuO remaining tightly in the cell wall even after extensive washing (Fig. 5). Synergistically, the layered Zn–CuO@GO could then wrap onto the bacteria surface to deprive of oxygen and nutrient exchange (II), owing to the layered structure of GO nanosheets, which led to a second disastrous inactivation.

To reveal the vivid mechanism of Zn–CuO@GO towards bacterial-killing as proposed in Scheme 1(II), we investigated their interactions with MDR *E. coli* and MRSA by TEM. Compared with Zn–CuO (ESI, Fig. S5†), the red arrow in Fig. 6 indicated the existence of GO sheets which was layer structured and well in combination with prickly Zn–CuO. As shown in Fig. 6A and B, the Zn–CuO@GO could bind to bacterial surface extensively due to their strong nonspecific interaction upon contacting, and distributed well throughout the cell wall (corresponding magnified figures were shown in Fig. 6a and b). To make it clearer, we also obtained zoomed figures illustrating the interaction between Zn–CuO@GO and MDR *E. coli* (Fig. 6), which clearly revealed that Zn–CuO@GO could then anchor and effectively penetrate into cellular membrane, to cause the cytoplasm leakage (cell lysis) subsequently. Especially, the Zn–CuO@GO also form a sublayer wrapping MDR *E. coli*, as indicated in Fig. 7A. It was also evident in Fig. 7B, that the anchored

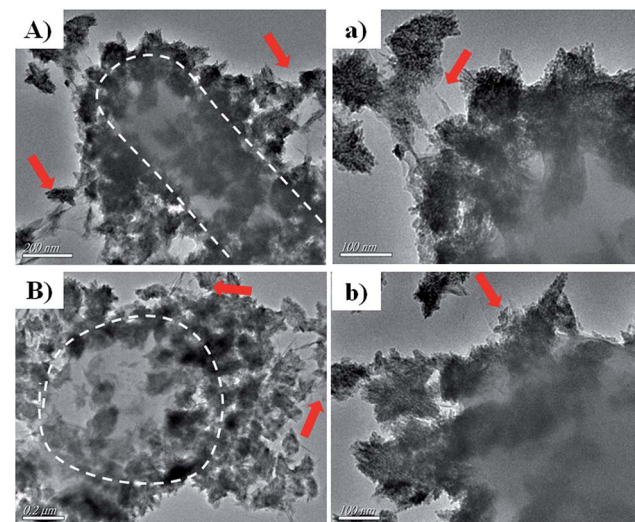


Fig. 6 (A and B) were TEM images of MDR *E. coli* and MRSA after treatment with Zn–CuO@GO, respectively; (a and b) were their corresponding magnified TEM images, respectively. White dashes outlined the borders of MDR *E. coli* and MRSA, and red arrows indicated the existence of GO as a sublayer for Zn–CuO@GO nanosheets.

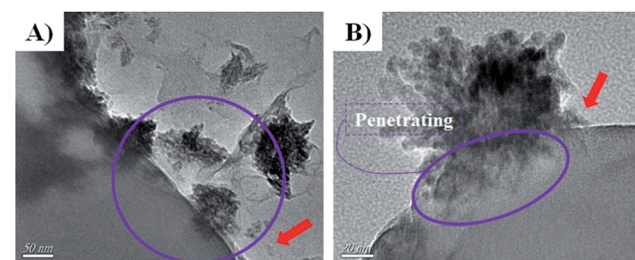
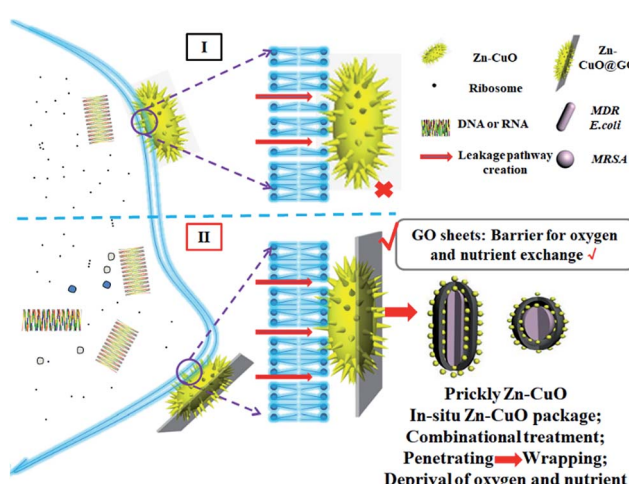


Fig. 7 (A and B) were further magnified TEM images uncovering the detailed antibacterial route of Zn–CuO@GO in association with MDR *E. coli* as an example.



Scheme 1 Schematic illustration of the bacterial killing mechanism of Zn–CuO@GO in comparison with prickly Zn–CuO.

Zn–CuO could exert their pierce effectively into the cell membrane. To this aspect, we concluded that Zn–CuO@GO could strongly bind to bacterial by wrapping the whole cell wall and dissociate the cell integrity, which also accounted for bacterial accumulation as indicated in Fig. S6 (ESI†), and verified the results obtained in Fig. 4. Thus, it is believable that the Zn–CuO@GO cause bacterial lysis from stepwise penetrating to wrapping upon contacting, namely, through exerting nano-pierce to damage cell walls and subsequently behave to wrap into the cell, which may deprive of oxygen and nutrient exchange and lead to ever-enhanced bacterial inactivation.<sup>7e</sup>

## Conclusions

In summary, we described a facile, bottom up and environment-friendly protocol for the scalable fabrication of Zn–CuO@GO nanocomposite for combinational bacterial-killing, especially towards multi-drug resistant strains, such as MDR *E. coli* and MRSA. Owing to their combinational antibacterial ability, the Zn–CuO@GO could accelerate bacteria lysis and inactivate

them nearly completely within 10 min upon contacting. Functioning in a stepwise penetrating and wrapping manner, the deposited prickly Zn-CuO in association with GO layer could penetrate and wrap into bacteria membrane to damage bacteria integrity, induce cytoplasm leakage, deprive of possible oxygen/nutrient exchange, and lead to progressive bacteria lysis. We thus concluded that suspensions containing Zn-CuO@GO nanosheets were promising disinfectant for combating multi-drug resistant bacterial stains from ICU to avoid severe bacterial infections.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The acknowledgements come at the end of an article after the conclusions and before the notes and references.

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