The neglected nano-specific toxicity of ZnO nanoparticles in the yeast *Saccharomyces cerevisiae*

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Nanoparticles (NPs) with unique physicochemical properties induce nano-specific (excess) toxicity in organisms compared with their bulk counterparts. Evaluation and consideration of nano-specific toxicity are meaningful for the safe design and environmental risk assessment of NPs. However, ZnO NPs have been reported to lack excess toxicity for diverse organisms. In the present study, the nano-specific toxicity of ZnO NPs was evaluated in the yeast *Saccharomyces cerevisiae*. Nano-specific toxicity of ZnO NPs was not observed in the wild type yeast. However, the ZnO NPs induced very similar nano-specific toxicities in the three mutants with comparable log $T_e$ (particle) values (0.64 vs 0.65 vs 0.62), suggesting that the mutants were more sensitive and specific for the NPs’ nano-specific toxicity. The toxic effects in the yeast were slightly attributable to dissolved zinc ions from the ZnO (nano or bulk) particles. Oxidative damage and mechanical damage contributed to the toxic effect of the ZnO particles. The mechanism of mechanical damage is proposed to be an inherent characteristic underlying the nano-specific toxicity in the mutants. The log $T_e$ (particle) was a useful parameter for evaluation of NPs nano-specific toxicity, whereas log $T_e$ (ion) efficiently determined the NPs toxicity associated with released ions.

Engineered nanomaterials (ENMs) with unique and novel physicochemical properties provide not only a lot of benefits but also serious threats once they enter into the environment. Theoretically, the special properties of ENMs can elicit nano-specific (excess) toxicity in organisms compared with their bulk counterparts1–3. Due to the potentially excess hazardous effect, the evaluation of the health safety and environmental risk of ENMs is becoming important and urgent. Fortunately, scientists have made great efforts in the safe design and risk assessment of ENMs. For instance, high content screening (HCS) had been successfully utilized to evaluate the environmental and toxicological impacts of NPs4. Although the biological and toxic mechanisms that induce toxicity in organisms have been extensively investigated5,6, nano-specific toxicities have been neglected in health risk assessments for various reasons. A large number of investigations confirmed the existence of nano-specific toxicities. For example, CuO nanoparticles (NPs) showed approximately 50-fold higher toxicity in *Vibrio fischeri*, *Daphnia magna* and *Thamnocephalus platyurus* than bulk CuO, and TiO2 NPs were shown to be toxic in *Daphnia magna* and zebrafish even though bulk TiO2 was not toxic3,7. A comparison of the toxicity of NPs and bulk particles suggested that a novel risk assessment pattern for NPs should be established, because the nano-specific effects are dramatically important and need to be seriously considered in the environmental assessment.

Interestingly, ZnO NPs exerted toxic effects analogous to ZnO bulk particles in various organisms, including bacteria (30 min EC50 in *Vibrio fischeri*, 1.9 vs 1.8 mg/L)3, algae (72 h EC50 in *Pseudokirchneriella subcapitata*, 0.042 vs 0.037 mg/L)7, yeast (8 h EC50 in *Saccharomyces cerevisiae*, 1.21 vs 1.34 mg/L)6, protozoa (24 h EC50 in *Thamnocephalus platyurus*, 0.18 vs 0.24 mg/L, and *Tetraphymena thermophila*, 6.8 vs 7.4 mg/L)3, crustaceans (48 h LC50 in *Daphnia magna*, 3.2 vs 8.8 mg/L)3, and fish (96 h LC50 in zebrafish, 4.92 vs 3.31 mg/L)7. These phenomena revealed that ZnO NPs lacked nano-specific toxicity, which conflicted with our general understanding that nanoparticles represented more serious hazardous threats to living organisms than bulk particles. Based on these results, whether ZnO NPs represent a similar hazard to the environment as bulk ZnO and whether the toxicological assessment pattern applied for bulk ZnO is also suitable for nano ZnO are unknown.

A previous study reported the nanotoxities of metal and/or metal oxide NPs derived from either NPs themselves, released metal ions, or both. For ZnO NPs, the released Zn2+ was proposed to be predominantly

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Table 1. Toxicity of nano ZnO, bulk ZnO and ZnSO4·7H2O for the yeast and three mutants based on 50% cell viability with the decreasing EC50. aEC50 of ZnSO4·7H2O to 4Δ was higher than 10129.00 mg/L due to the high concentration of zinc salt that induced 6.54 (%) cell death. b Toxicity differences between nano ZnO and bulk ZnO calculated through equation 3. c Td (particle) and Td (ion) denote the toxic ratio of bulk ZnO or zinc ion toxicity to ZnO NP toxicity calculated by equations 4 and 5, respectively.

| Yeast and mutants | EC50 (mg/L) | Td (particle) | Td (ion) | log Td (particle) | log Td (ion) |
|-------------------|-------------|---------------|----------|------------------|-------------|
|                   | Nano ZnO    | Bulk ZnO      | ZnSO4·7H2O |                  |             |
| BY4741            | 8.83 (±0.52) | 11.66 (±1.18) | 498.06 (±68.43) | 2.83            | 1.32        |
| yap1Δ             | 12.67 (±1.07) | 55.82 (±6.34) | 227.07 (±23.37) | 43.15          | 4.41        |
| 4Δ                | 6.02 (±0.32)  | 26.72 (±2.91) | >101290.00a | 20.70          | 4.44        |
| 5Δ                | 4.34 (±0.39)  | 18.09 (±1.47) | 281.34 (±25.34) | 13.75          | 4.17        |

The toxicity was significantly more serious compared to other nano or bulk ZnO particles only when the NPs size was shorter than the dissolved Zn2+ or ZnO (nano and bulk) Particles and Zinc Salt Toxicity towards Saccharomyces cerevisiae. According to the investigation of the predominant difference between nano and bulk ZnO is the particle size. According to the investigation of the size-dependent toxicity of ZnO NPs, the ZnO NPs toxicity was not governed by particle size or its related factors. The toxicity was significantly more serious compared to other nano or bulk ZnO particles only when the NPs size was less than 20 nm. The possibility of aggregation in the present investigation suggested that the ZnO NPs toxicity was not dependent on the particle size, which was reported to strongly affect Zn2+ release. Thus, the dissolved Zn2+ could not be strongly correlated with the toxicity (as detailed below). Notably, in the present study some of the ZnO bulk particle sizes still belonged to the nanoscale (<100 nm, see Fig. S1).
Herein, the mutant strains exhibited varied toxicity patterns in BY4741. Increased differences in toxicity were observed when the nano ZnO was compared with bulk ZnO (yap1Δ: $T_d = 43.15$ mg/L; 4Δ: $T_d = 20.70$ mg/L and 5Δ: $T_d = 13.75$ mg/L). We hypothesized that these increased toxicity differences were ZnO NPs nano-specific. Previously, single gene deletion yeast mutants were used to examine the nanotoxicity of CuO NPs. The yap1Δ mutant showed higher tolerance to CuO NPs toxicity15,16. This result was consistent with our result that nano and bulk ZnO yielded lower toxicity in yap1Δ, which was reflected by the higher EC$_{50}$ compared to the wild type. Moreover, yap1Δ showed similar tolerance to other NPs, such as Ag NPs and Ag$_2$O NPs (data not shown). Theoretically, yap1Δ should be sensitive to ROS-mediated toxicants. Indeed, intercellular ROS investigations confirmed that the yap1Δ strain was sensitive to ZnO NPs (see below), and CuO NPs induced more ROS than the wild type strain16. The comparative toxicities of these NPs in yap1Δ and the wild type strain could possibly be explained by their toxic mechanisms, which slightly accounted for ROS generation. However, the rigid cell wall of the yeast can prevent the nano or bulk particles from entering the intercellular space and stop the triggering of the deficient defense system that is hypersensitive to oxides such as hydrogen peroxide. Thus, in this instance, the observed toxicity is only governed by mechanical damage, such as that described here. Alternatively, the reported compensatory mechanism in the cell may mask the deficiency of a single gene's desired phenotype20,21.

Correspondingly, multiple gene deletion mutants can be more sensitive and adequate for cytotoxicity estimations. As expected, the quadruple (4Δ) and quintuple (5Δ) mutants were more sensitive than the wild type and yap1Δ strains to ZnO NPs with lower EC$_{50}$ values (6.02 and 4.34 mg/L, respectively). A similar phenomenon was observed for CuO NPs16, suggesting the sensitivity and specificity of these mutant strains to nanotoxicity. Considering the deficiencies in the 4Δ and 5Δ gene mutants20,21, the observed serious toxic effects could be assumed to correlate with ROS and/or cell membrane permeability. Furthermore, although they presented decreased toxicity differences (26.70 and 13.75 mg/L, respectively), their nano-specific toxicities were still detectable (log $T_d$ (particle) ≈ 0.65). Interestingly, lower toxic effects were observed for the 4Δ and 5Δ mutants during contact with bulk ZnO compared with the wild type yeast. However, further investigations are needed to understand the phenomenon.

Mechanisms underlying the nano-specific toxicity of ZnO NPs. Three major mechanisms were previously proposed to be involved in the toxic actions of metals and metal oxide NPs5,6,22: (I) Released metal ions from particles (e.g., Zn$^{2+}$ from ZnO NPs); (II) ROS generated from interactions between NPs and biological targets (e.g., -OH or O$_{2•}^{-}$); and (III) mechanical damage due to the direct interactions of NPs with biological targets (e.g., broken cell membranes). Herein, the three toxic mechanisms were examined and detailed below individually to elucidate the mechanism by which ZnO NPs induce nano-specific toxicity in organisms.

**Dissolved Zn$^{2+}$.** Recently, many scientists reported that the mechanism underlying the toxicity of ZnO NPs was dependent on the release of Zn$^{2+}$. Indeed, many investigations found that ZnO NPs displayed toxicity in organisms comparable to those targeted with bulk ZnO and zinc salts (ZnCl$_2$ or ZnSO$_4·7$ H$_2$O). Nevertheless, as noted previously the Zn$^{2+}$ concentrations released from ZnO NPs were too low to explain the toxic injury9,23, indicating that additional mechanisms (i.e., solid ZnO NPs) played a role in the toxic effect.

As shown in Fig. 1, the Zn$^{2+}$ release curves for nano and bulk ZnO were very close at the 1 and 10 mg/L concentrations, although more Zn$^{2+}$ was released from nano ZnO than bulk ZnO at the highest concentration (100 mg/L). At the 1 mg/L concentration, nano ZnO dissolved rapidly and completely. This finding was confirmed by the Noyes-Whitney equation24. In contrast, less than 10% of the nano and bulk ZnO were dissolved at the primary concentrations of 10 and 100 mg/L, which was consistent with previous reports8,10. Both nano and bulk ZnO illustrated stronger toxic effects in the set of yeasts than the zinc salt (ZnSO$_4·7$ H$_2$O) (see Fig. S2 or Table 1). Thus, the Zn$^{2+}$ released from either the ZnO NPs or bulk ZnO was not sufficient to elicit serious toxicity. Even when a high concentration of Zn$^{2+}$ was employed (101,290.00 mg/L), a less than 10% decrease in cell viability was detected in the quadruple mutant (4Δ). The more than two log unit higher toxicity of the nano and bulk ZnO strongly indicated that the mechanism underlying the toxicity in yeast was essentially dependent on
the particles themselves and not on the released zinc ions. However, we evaluated the extent to which the toxic effect was attributable to the dissolved zinc ions using previously described methods. As shown in Fig. S3, the calculated concentrations of nano and bulk ZnO led to an almost 50% decrease in cell viability, with EC50 ranges from 44.16 to 57.35%. This result indicated that the cell viability testing was valid. Moreover, Zn2+ released from either nano ZnO or bulk ZnO yielded less than 9% cell death (range 3.16 to 8.25%, see Fig. S3). Correspondingly, the contributions of Zn2+ to the overall toxic effect accounted for less than 8% (range 2.93 to 7.44%; see Table 2). Zn2+ sometimes did not induce toxic effects in the yeast or mutants, resulting in the larger error bars in Fig. S3. Moreover, the investigation of intercellular ROS demonstrated that dissolved Zn2+ did not induce significant oxidative stress in the yeast (p > 0.05) compared to the ZnO particles (see Fig. 2). Because the solubilities of metals and metal oxides vary and most metal ions induce diverse toxic effects in organisms, the concentrations of metal ions released from NPs should be routinely measured and provided in the ecotoxicological and environmental assessments.

Additionally, the intercellular Zn2+ could be another toxic source. Theoretically, 4Δ mutant with cell membrane permeability defect should be more sensitive to free ion-related toxic effect than BY1437, because the released Zn2+ could be inside of 4Δ mutant freedom. Therefore, the concentration of intercellular Zn2+ should be higher in 4Δ mutant cell than in BY1437 cell. Once the intercellular Zn2+ can induce serious toxic effect, the applied lower concentration of zinc salt should be inducing serious toxic effect in 4Δ mutant. However, when a high concentration of Zn2+ was employed (101,290.00 mg/L), a less than 10% decrease in cell viability was detected in the 4Δ. Furthermore, the applied comparable free Zn2+ (ZnSO4·7H2O) did not induce severe intercellular oxidative stress (p > 0.05) in the yeast strain. Comparably, the ZnO NPs caused significant intercellular oxidative stress (p < 0.05 or 0.01). The combination of these results revealed that the toxic effect of nano or bulk ZnO was not mainly dependent on the released Zn2+ but on particle-related toxicity (i.e., oxidative stress and/or mechanical damage). Importantly, the particles contributed to the comparable toxic effect in the set of yeasts (almost 95%, see Table 2); hence, the dissolved Zn2+ was not the essential distinction for understanding why the ZnO NPs did not induce higher toxicity than bulk ZnO.

**Intercellular ROS.** In addition to the dissolved metal ions, ROS generation was proposed to be a general paradigm for the toxic action of metal and metal oxide NPs. The induced intercellular ROS could lead to dysfunctions in mitochondrial oxidative phosphorylation and ultimately cause cell death. Therefore, intercellular ROS generation was explored to evaluate whether oxidative stress was the major toxic mechanism of ZnO particles and the inherent factor in nano-specific toxicity elicitation. Our previous investigation demonstrated that the multiple gene-deleted mutants were sensitive to NPs-generated oxidative stress. As shown in Fig. 2, the selected highest concentration (2 mg/L) of ZnO particles generated significantly (compared to wild type) and extremely...
significantly (compared to the other mutants) higher ROS than the control group ($p < 0.05$ and $p < 0.01$). Comparably, $0.1 \text{ mg/L of ZnO particles did not induce serious oxidative stress in the yeast, although significant oxidative stress was detected in the yap1Δ mutant following exposure to 0.1 mg/L of ZnO NPs (p < 0.05, see Fig. 2B). Oxidative stress induced by these yeast strains was aggravated by increasing concentrations of ZnO nanoparticles, suggesting that oxidative damage at least partially contributed to the toxic effect. Similarly, ZnO NPs exerted antibacterial activity against S. cerevisiae via the induction of oxidative stress$^{12}$.

Interestingly, nano and bulk ZnO showed similar patterns of oxidative stress in the yeast mutants, with either no extraordinary oxidative stress at the 0.1 mg/L concentration (expect for yap1Δ) or (extremely) significantly high oxidative stress at the 2 mg/L concentration. The lack of significant differences in the effects of the ZnO particles on the yeast strains ($p > 0.05$) suggested that oxidative stress was not an essential characteristic to differentiate ZnO NPs nano-specific toxicity.

Notably, extracellular ROS generated in response to the interaction of the particles with cellular membranes (walls) also contributed to the antibacterial activity of ZnO NPs$^{26,27}$. As reported earlier, particle-dependent ROS formation perturbed electron transfer processes in the cell, such as those in the mitochondrial inner membrane$^{23}$. Extracellularly generated ROS can also lead to cell membrane damage, cell lysis and ultimately cell death$^{28,29}$. Theoretically, the primary location for extracellular ROS generation is the yeast cell wall. This conclusion can be confirmed by the data of zeta potentials of the particles and yeast strains (see below). Hence, the generated ROS from solid ZnO on the surface of the yeast walls could be an additional toxic source in this study. The extracellular ROS could possibly serve as helper to destroy the cell walls or membranes, and then enter into the internal yeast cell. Indeed, Fig. S4 suggested that the yeast cell wall and membranes were partly destroyed. Although without of ZnO NPs adhering to the cell wall or membrane was found through TEM, extracellular ROS might be generated at the extracellular wall NPs. If ZnO NPs are adsorbed on the surface of the yeast cell walls, but the microenvironment could promote dissolution of ZnO NPs on the cell wall, resulting shift of particle-related toxic effect to ion-related toxic effect. Therefore, the ROS or other relatively biological mechanisms could attribute to the cytotoxicity, although the possibly toxic effects were not quantitatively measured in this study.

Mechanical damage. The TEM images indicated that the yeast morphologies following ZnO NPs exposure were dramatically different compared with normal yeast cells (see Fig. 3). The cell walls of some of the yeast were broken or partially broken (Fig. 3C,D). The phenomenon was also observed in E. coli$^{23}$. In the other yeast, although the cell walls were not crushed completely, the ZnO NPs-treated cell walls were deformed with some sunken areas (Fig. 3F) or deficiencies (Fig. 3E). Figure 3E indicated that the yeast cell walls were disrupted and that the morphology of the normal cell changed from globular to an irregular shape.

Regardless, the rigid yeast cell wall could protect the yeast cells from ZnO NPs-induced mechanical damage. Moreover, as shown in Fig. S4, the yeast cell wall prevented the passage of the Annexin V-FITC dye (Q4 region) but not PI (Q1 region), suggesting that most of the rigid cell walls were not crushed. This cell wall protection was also reported for several other organisms, such as S. typhimurium, K. pneumonia, and S. aureus$^{28}$, in which breakage of the cell wall or membrane was not detected. Correspondingly, these sunken areas suggested that endocytosis could be a potential route for the entrance of ZnO NPs into the cell. However, the cell membrane was fragmented under the sunken areas as indicated with the red arrow in Fig. 3D,F. The disorganization or permeability of the yeast membrane during nano and/or bulk ZnO exposure can lead to the loss of cell cytoplasm and ultimately to cell death. As shown in Fig. 3B, the cell cytoplasm leaked and passed through the cell membrane, although the rigid cell wall prevented the cytoplasm from leaking out. Furthermore, without of intercellular ZnO NPs into cell or adhesion of ZnO particles on cell wall or membrane were detected by TEM in this study. These findings suggested that the ZnO particles could not be taken into by the yeast strains. Comparably, ZnO NPs exposed led to Escherichia coli cell membrane fragmentation and deformation$^{10,30}$. Without the protection of the tough cell wall, ZnO NPs were detected in the inner bacterial cell$^{29–31}$. Furthermore, ZnO NPs arrived in certain fish organs through waterborne routes or ingestion and subsequently caused tissue damage$^{28,32}$. For example, ZnO NPs induced shrinkage and loss of the cell cytoplasm in zebrafish gills$^{2}$, hyperplasia of the gall filament epithelium and edema of the carp gill lamellae$^{31}$. ZnO NPs were also observed to alter the structural and functional integrity of cell junctions in the Xenopus laevis intestine$^{22}$. Despite the dissolved Zn$^{2+}$ was not shown to induce serious cytotoxicity here, the “Trojan horse effect” derived from the internalized NPs was seriously considered$^{28,35}$. Although we examined the yeast cells very carefully by TEM, no intercellular ZnO particles were found. This result suggested that the ZnO particles could not be taken into by the yeast strains. Interestingly, our recent study found similar phenomenon inside yeast cells using CuO NPs that no CuO NPs was detected by TEM. Even if a very little amount of ZnO particles could enter the yeast cell, they should rapidly dissolve in the acidic environment of the lysosome (pH 5). Based on the combination of qualitative inspection by TEM and the quantitative data presented in Fig. S4 (Q4 region), the rigid cell wall of the yeast was barely broken and the ZnO particles scarcely entered the intercellular area of the yeast cells. Therefore, the “Trojan horse effect” was not validated to yeast strain in this study. Theoretically, determining the uptake routes of ZnO NPs into organisms with or without of cell walls are very important for the toxicity inducement and risk assessment.

Undoubtedly, the mechanical damages caused by particles binding to the cell surface are majorly governed by electrostatic forces, and hence the zeta potentials of the particles and yeast strains were measured (see Table S1). All yeast strains showed the expected negative charge, and all of their negative charges were slightly decreased at the 6 h time point. Interestingly, the ZnO NPs possessed a positive charge that was absolutely different from the negative charge of the bulk ZnO. The yeast strains preferred to bind to the ZnO NPs but not to the bulk ZnO. Theoretically, more serious cell wall and/or membrane damage would emerge under ZnO NPs exposure compared to bulk ZnO. Indeed, this finding was confirmed by the PI dye experiment shown in Fig. 4, in which ZnO NPs at concentrations of 5 and 10 mg/L induced stronger cell permeability in yap1Δ, 4Δ and 5Δ compared to bulk ZnO.
Interestingly, no significant difference in cell permeability was detected between the nano and bulk ZnO in the wild type strain (p > 0.5, see Fig. 4A). One possible explanation was that the mechanical disturbance under the shaking condition (200 rpm) overcame the electrostatic forces and contacts that existed between the bulk ZnO and the cell wall.

Massive ROS species generation could also lead to cell membrane damage and ultimately cell death. In the present investigation, intercellular ROS generation could not possibly lead to cell membrane damage because the

Figure 3. The morphology of yeast strain cells exposed to nano ZnO (10 mg/L). (A) normal cell, (B) cell cytoplasm leakage, (C) crushed cell wall, (D) partially crushed cell wall, (E) irregularly shaped cell, (F) deformed cell with sunken area.
relative ROS concentrations generated during exposure to ZnO NPs and bulk ZnO were comparable \( (p > 0.05) \) in each yeast strain. In contrast, the mechanical damage showed different patterns between the wild type and mutants that could primarily account for their various physiologies. The deletion of a gene or genes affected the cell defense mechanism, which was very sensitive and specific for nanotoxicity. In the wild type yeast, the sensitivity or specificity could be masked by an influencing factor, such as the shaking described here.

In summary, the free Zn\(^{2+}\) released from ZnO (nano and bulk) particles were just slightly contributed to the detected cytotoxicity. Oxidative damage, possibly from both extra- and inter-cellular ROS, partially contributed to the toxic effect of the ZnO particles. Additionally, mechanical damage was another mechanism underlying the toxicity of ZnO NPs and most likely accounted for the nano-specific toxicity of ZnO NPs in the mutants.

Toxicity Differences and Toxicity Enhancement of ZnO NPs in the Wild Type and Mutants.

As summarized in Table 1, the ZnO particles displayed similar cytotoxicity in the wild type yeast \( (E_{50} = 8.83 \text{ (nano)} vs 11.66 \text{ (bulk)} \mu g/L; \; T_d = 2.83 \mu g/L) \). However, different cytotoxicity patterns between bulk ZnO and nano ZnO were detected in the mutants \( (T_d = 43.15 \mu g/L \text{ for yap1Δ}; \; T_d = 26.70 \mu g/L \text{ for 4Δ and } T_d = 13.75 \mu g/L \text{ for 5Δ}) \). Therefore, the nano-specific toxicities of ZnO NPs in the yeast were not exhibited towards the wild type but were exhibited towards the three mutants. There was a decreasing tendency for nano-specific toxicity in BY1437 < 5Δ < 4Δ < yap1Δ. However, the tendency conflicted with the toxicity results, where 5Δ presented the highest sensitivity to ZnO NPs, followed by 4Δ, BY 1437 and yap1Δ. Indeed, we previously observed that yap1Δ was less sensitive to NPs toxicity, and the ZnO NPs led to the lowest cytotoxicity for yap1Δ herein (see Table 1). The strongest cytotoxicity of the ZnO NPs was detected for the 5Δ strain, which exhibited almost two-fold higher toxicity than the wild type. These observed toxicity differences were due to the strains’ distinct physiologies and related toxic mechanisms (as discussed above). Compared to the wild type, the three mutants lacked sensitivity to bulk ZnO, which had a higher \( E_{50} \) (11.656 vs 55.82 vs 26.72 vs 18.09 \( \mu g/L \)).

In parallel with the toxicity differences, toxicity enhancement \( (T_e) \) is another important parameter to evaluate the nano-specific toxicity of NPs. From the organic chemistry toxicity perspective, toxicity enhancement \( (T_e) \) had been successfully utilized to discriminate and evaluate the excess toxicity of organic chemicals\(^{37}\). \( T_e = 10 \) was identified as a threshold to separate excess toxicity \( (\log T_e \geq 1) \) from baseline toxicity \( (\log T_e < 1) \). Theoretically, nanoparticles should exert seriously harmful damage compared to their bulk counterparts with identical chemistry because the toxicity is nano-specific. Thus, the nano-specificity of the NPs toxicity could be expressed as the toxicity enhancement. In the present investigation, a hypothesis was proposed that bulk particles only exerted baseline toxicity but NPs possibly exhibited excess toxicity due to their unique physicochemical properties. Therefore, it is likely that the nano-specific toxicity of NPs in the target organisms can be expressed as excess toxicity compared with the bulk particles.
ZnO NPs and zinc salt were identical. The log of dissolved Zn\(^{2+}\) compared with Cu\(^{2+}\) indicated that the mechanism underlying the toxicity of the ZnO NPs in yeast was not dependent on dissolved Zn\(^{2+}\). Our findings were consistent with previous reports showing that the ZnO NPs induced toxicity in yeast that was not dependent on dissolved metal ions 

**Table S2.** Relative toxicity contribution of particles and released ions from nano or bulk ZnO to the nominal EC\(_{50}\) in the yeast and three mutants.

| Yeast strains | Particle contribution (%) from ZnO NPs | Released ion contribution (%) from Zn\(^{2+}\) | Bulk ZnO | Bulk CuO |
|---------------|----------------------------------------|---------------------------------------------|---------|---------|
| BY4741        | 93.36                                  | 6.64                                        | 7.44    |         |
| pop1Δ        | 95.72                                  | 4.28                                        | 4.12    |         |
| Δ             | 96.68                                  | 3.32                                        | 4.58    |         |
| ΔΔ            | 95.98                                  | 4.02                                        | 2.93    |         |

As shown in Table 1, the log \(T_c \) (particle) for the wild type yeast was 0.12, indicating that no excess toxicity was found. Thus, the ZnO NPs exhibited no nano-specific toxicity for the wild type yeast. Not only was this phenomenon previously found in wild type yeast but similar results were also reported for other organisms. As shown in Table S2, the log \(T_c \) (particle) of the ZnO NPs was less than 0.5, suggesting that the ZnO NPs induced a level of toxicity similar to bulk ZnO but did not induce nano-specific toxicity. In contrast, CuO NPs are an excellent example of NPs that induce nano-specific toxicity in target organisms (see Table S2). Regardless of the target organisms, toxic endpoints or exposure duration, log \(T_c \) (particle) values of CuO NPs were higher than those of ZnO NPs. Interestingly, all three mutant strains possessed dramatically similar log \(T_c \) (particle) values (0.64 \(\pm\) 0.05 vs. 0.65 \(\pm\) 0.10). Similar to the \(T_c \) (particle) values, the log \(T_c \) (particle) showed that the ZnO NPs exerted at least partial nano-specific toxicity for the mutant strains, although the log \(T_c \) (particle) was less than 1. Importantly, the ZnO NPs yielded almost identical nano-specific toxicities in the mutant strains, revealing that the three mutants possessed almost identical sensitivity and specificity for the ZnO NPs nano-specific toxicity. \(T_c \) (particle) can exactly reflect the excess NPs toxicity in terms of the toxicity difference, although the latter parameter considers the specificity of the physiology of diverse strains.

Different from \(T_c \) (particle), \(T_c \) (ion) exhibited a varied pattern of excess toxicity in the wild type yeast and mutants. The log \(T_c \) (ion) covered almost 3 log units (from 1.35 (BY) to 4.23 (ΔΔ)), suggesting that ZnO NPs induced extremely excessive toxicity compared to Zn\(^{2+}\). Moreover, the log \(T_c \) (ion) was greater than 1, indicating that the mechanism underlying the toxicity of the ZnO NPs was not dependent on dissolved Zn\(^{2+}\). As shown in Table S2, different results suggested that the ZnO NPs exhibited toxicity that was comparable with Zn\(^{2+}\). Indeed, the log \(T_c \) (ion) equaled zero for T. thermophila and P. subcapitata, indicating that the toxicities of the ZnO NPs and zinc salt were identical. The log \(T_c \) (ion) in V. fischeri was negative 0.24, suggesting the Zn\(^{2+}\) showed slightly higher toxicity than the ZnO NPs. As mentioned above, the possible dissolved ratio of Zn\(^{2+}\) in the NP suspensions was not sufficient to induce toxicity to a level comparable with the zinc salt. Thus, it was an oversimplification to assert that the mechanism underlying the ZnO NP toxicity was primarily dependent on dissolved Zn\(^{2+}\) based on the similar toxic effect observed for the ZnO NPs and zinc salt. Enrique Navarro et al. corrected this shortcoming by examining the functions of metal ion concentrations. Importantly, log \(T_c \) (ion) of Ag NPs (from 1.35 to 4.23 (4Δ)) confirmed the finding that the released Ag\(^{+}\) was a major toxicity resource, although Ag\(^{+}\) could not fully explain the toxicity in Chlamydomonas reinhardtii. Moreover, based on the inhibition in E. coli following exposure, a log \(T_c \) (ion) \(< -1\) for Ag NPs was proposed that supported the reported result that the toxicity was directly correlated with the released metal ions. Furthermore, comparable log \(T_c \) (ion) values (−0.89 vs. −0.91, very close to −1.00) were calculated for two differently coated Ag NPs in Eisenia fetida, thereby confirming that the proposed mechanism of toxicity was dependent on the released Ag\(^{+}\).

The CuO NPs exhibited varied toxicity patterns compared with the ZnO NPs. With the exception of S. cerevisiae, the log \(T_c \) (ion) values of the CuO NPs were less than negative 1.30, revealing that Cu\(^{2+}\) showed more than 10-fold higher toxicity than the CuO NPs. The different log \(T_c \) (ion) values of the ZnO and CuO NPs strongly indicated that the mechanisms underlying the ZnO NPs toxicity were not associated with the released Zn\(^{2+}\), whereas the mechanism underlying the CuO NPs toxicity was associated with the released Cu\(^{2+}\). Notably, differences in the log \(T_c \) (ion) values between the ZnO and CuO NPs could also occur because Zn\(^{2+}\) showed less toxicity compared with Cu\(^{2+}\). The Zn\(^{2+}\) and Cu\(^{2+}\) concentrations that induced growth toxicity ranged from 200 to 300 and 15 to 20 \(\mu\)g/g dry weight, respectively. Moreover, Cu\(^{2+}\) formed more stable complexes with proteins and/or peptides than Zn\(^{2+}\), suggesting that the Cu\(^{2+}\)-mediated toxicity was more irreversible and serious. In S. cerevisiae, the log \(T_c \) (ion) of the CuO NPs was an outlier at 0.12, indicating that the toxicity of the CuO NPs almost equaled the toxicity of Cu\(^{2+}\). This result was most likely caused by the testing circumstances: growth medium (log \(T_c \) (ion) = −0.10, growth inhibition) vs deionized water (log \(T_c \) (ion) = −0.77, cell viability). Based on our unpublished cell viability data, the log \(T_c \) (ion) of CuO NPs in deionized water is −0.97, suggesting that the effect of Cu\(^{2+}\) on cell viability is almost 10-fold higher than the effect of CuO NPs. As mentioned above, proteins in the growth medium could strongly capture Cu\(^{2+}\) and hence decrease the toxicity of Cu\(^{2+}\).

These findings suggested that the mechanism underlying the ZnO NPs toxicity was mainly related to the released Zn\(^{2+}\). The present results revealed that log \(T_c \) (particle) could be a useful parameter to evaluate whether certain NPs induce nano-specific toxicity in target organisms, whereas log \(T_c \) (ion) could be employed to understand whether the mechanism of the toxicity of the metal or metal oxide NPs correlated with the corresponding dissolved metal ions.

**Conclusions**

In summary, ZnO NPs displayed nano-specific toxicity in the yeast mutants, although no nano-specific toxicity was observed in the wild type yeast. Oxidative damage partially contributed to the adverse effects in the yeast.
strains but did not contribute to the nano-specific toxicity of the ZnO NPs in the mutants. Mechanical damage also attributed to the adverse effects of the ZnO NPs in the yeast strains and was the essential mechanism underlying the nano-specific toxicity in the mutants. The nano-specific toxicity of the ZnO NPs' could potentially lead to serious environmental hazards and threaten human health; thus, the underestimated nano-specific toxicity should be carefully considered in the risk assessment. In particular, log $T_e^{(\text{particle})}$ is a useful parameter to estimate the NPs’ nano-specific toxicity in target organism, whereas log $T_e^{(\text{ion})}$ is an efficient parameter to determine whether the NPs’ toxicity mechanism is primarily correlated with its released ions.

Methods

Test organisms and chemicals. The basic information for the yeast S. cerevisiae strains is summarized in Table S1. The cultivation conditions and procedures for the S. cerevisiae strains were detailed previously56. The viability of the yeast strains and the effect of the pH value during the growth phase were explored previously16 and were shown in Figs S5 and S6. The cell viabilities of these yeast strains under non-stress conditions at different time points were measured and displayed in Fig. S7. The combination of the above information and the cell viabilities of selected strains were not significantly different under non-stress conditions, and hence the detected cytotoxicity was due to the employed chemicals.

ZnO NPs (20 nm) were purchased from Beijing Nachen S &T Ltd.; the purity was 99.9%. Bulk ZnO and ZnSO$_4$$\cdot$$7$H$_2$O, with over 99.9% purity were purchased from Tianjin Kemiou Chemical Reagent Co. Ltd., China. Considering the great purity of these materials, the toxic effects detected in this investigation were derived from the materials.

The shape and size of the ZnO particles in suspension were visualized by transmission electron microscopy (TEM, JEM-100CXII, JEOL, Ltd., Japan). The released Zn$^{2+}$ from three concentrations (1, 10 and 100 mg/L) of ZnO particles in the absence of light at room temperature was detected by Atomic Absorption Spectroscopy. The zeta potentials and hydrodynamic diameters of the ZnO particles and the zeta potentials of the yeast strains were characterized at the 1 and 6 h time points using a Nano-Zetasizer (1000HS, Malvern Instrument Ltd., UK).

Cytotoxicity test. Both nano and bulk particles were prepared at high concentrations in deionized water and then stored in the dark after ultrasonication for 30 min. Similarly, ZnSO$_4$$\cdot$$7$H$_2$O was employed for Zn$^{2+}$ stock solution preparation without ultrasonication.

The cytotoxicity test for yeast cell viability was performed as follows. Yeast in the logarithmic growth phase after overnight cultivation were harvested and washed three times with sterile deionized water. The yeast pellets were resuspended and diluted using sterile deionized water until the cell density (optical density, OD$_{600}$) was equal to 1.050 (±0.030). Nominal concentrations of the test compounds were obtained through the 10-fold dilution method. Under sterile conditions, at least six different concentrations (with three replications) and a blank control were evaluated. After six hours of exposure (200 rpm, 30 °C), 100 μL of the test suspension for each concentration was diluted and spread onto YPD solid plates. The colonies were counted after two days of growth and compared with the control.

The EC$_{50}$ values were calculated using dose-response cytotoxicity curves generated from a confirmed four-parameter sigmoid function equation (1)37:

$$y = \min + \frac{(\max - \min)}{1 + \left(\frac{x}{\text{Hillslope}}\right)^{\text{EC}_{50}}}$$

(1)

The calculated EC$_{50}$ values of the nano and bulk ZnO served to determine the relative contributions of the nano and bulk particles and their released zinc ions to the cytotoxicity of the yeast and mutants. Part of a suspension with a nominal EC$_{50}$ was utilized to examine the overall inhibition. The residual of the suspension was centrifuged (10,000 x g, 10 min) and then the supernatant was passed through a 0.22 μm filter to obtain the released Zn$^{2+}$. The released Zn$^{2+}$ was applied to explore the ions’ contribution to the cytotoxicity. The details of the toxicity exposure were the same as described above. The relative contributions to the cytotoxicity from the particles or ions were obtained through equation 2 based on the hypothesis that ZnO particles and Zn$^{2+}$ possessed different modes of action as reported previously11.

$$E_{(\text{total})} = 1 - \left[\left(1 - E_{(\text{ions})}\right)\left(1 - E_{(\text{particle})}\right)\right]$$

(2)

in which $E_{(\text{total})}$ indicates the total detected toxic effect derived from the suspensions, $E_{(\text{ions})}$ indicates the detected toxic effect from the dissolved Zn$^{2+}$, and $E_{(\text{particle})}$ denotes the nano or bulk ZnO particle-induced toxic effect.

Intracellular ROS. Intracellular ROS generation resulting from contact with the ZnO NPs, bulk ZnO and Zn$^{2+}$ was detected using the fluorescent probe 2′,7′-dichlorofluorescin diacetate (H$_2$DCFDA). After 6 hours of toxic exposure, the yeast cells were harvested and washed three times with 0.1 M PBS (pH = 7.20). The cell pellets were resuspended and 10 μM H$_2$DCFDA was added; then, the cells were incubated for 30 min in the dark. Subsequently, the cells were washed three times with PBS, and the cell pellets were suspended with a final biomass of 1–5 × 10$^5$ cells/ml to measure the fluorescence. Untreated yeast and H$_2$O$_2$ (10 μL, 30%) served as the control and positive control, respectively. The measurement results of the treatment groups were expressed as the relative ROS values compared with the control group.
Mechanical damage determination. To detect apoptotic cell death, an Annexin V-FITC/PI double dye kit (Nanjing Jiancheng Bioengineering Institute, China) was used. The results are exhibited in Fig. S4. Due to the cell wall protection, PI (propidium iodide) dye was used to evaluate cell membrane permeability. The PI was dissolved into 0.1 M PBS (pH = 7.20) to a final concentration of 50 μg/mL and added to the yeast with a final biomass of 1–5 × 10^5 cells/mL. Untreated yeast and H_2O_2 (10 μL, 30%) served as the control and positive control, respectively. High speed sorting flow cytometer (FACSAria™ III, DB Company, USA) was used to analyze the nano-specific toxicity compared with the bulk counterpart.

The cell wall protection, PI (propidium iodide) dye was used to evaluate cell membrane permeability. The PI was dissolved into 0.1 M PBS (pH = 7.20) to a final concentration of 50 μg/mL and added to the yeast with a final biomass of 1–5 × 10^5 cells/mL. Untreated yeast and H_2O_2 (10 μL, 30%) served as the control and positive control, respectively. High speed sorting flow cytometer (FACSAria™ III, DB Company, USA) was used to analyze the toxicity of the bulk ZnO over the toxicity of the ZnO NPs (see equation 4).

Toxicity difference and toxicity enhancement. Herein, the toxicity difference (T_d) was defined as the difference in the EC_{50} between the ZnO NPs and bulk ZnO for the yeast and its mutants. The calculation was performed using equation 3. Notably, the increasing toxicity of the particles decreased the EC_{50}.

\[ T_d = \text{EC}_{50} (\text{bulk particle}) - \text{EC}_{50} (\text{nano particle}) \]  

Toxicity enhancement (T_e^{particle}) was proposed and defined as the excess toxicity of NPs resulting from their nano-specific toxicity compared with the bulk counterpart. T_e^{particle} was a dimensionless toxicity ratio calculated by the toxicity of the bulk ZnO over the toxicity of the ZnO NPs (see equation 4).

\[ T_e^{particle} = \frac{\text{EC}_{50} (\text{BP})}{\text{EC}_{50} (\text{NPs})} \]  

T_e^{ion} was used to evaluate whether the toxic mechanism was dependent on metal ions. The comparison of ion toxicity (EC_{50} (ion)) and NPs (EC_{50} (NPs)) yielded T_e^{ion} in terms of toxicity difference.

\[ T_e^{ion} = \frac{\text{EC}_{50} (\text{ion})}{\text{EC}_{50} (\text{NPs})} \]  

Statistical analysis. All quantitative data were shown as the mean ± SD of a representative from at least three independent experiments. Triplicate tests were performed for each experiment. The t test was used to compare the differences between two groups. P < 0.05 (*) and P < 0.01 (**) were considered statistically significant and extremely statistically significant, respectively.

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Acknowledgements
We thank Yan Wang from the Technical Service Center of the Institute of Hydrobiology for providing technical support and gratefully acknowledge funding support from the National Natural Science Foundation of China (grants No. 21477159 and No. 21037004).

Author Contributions
WC.Zhang and T.F. designed the experiment(s). WC.Zhang and SP.Bao conducted the experiment(s) and analyzed the data. WC.Zhang wrote the manuscript, and Tao Fang refined the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Zhang, W. et al. The neglected nano-specific toxicity of ZnO nanoparticles in the yeast Saccharomyces cerevisiae. Sci. Rep. 6, 24839; doi: 10.1038/srep24839 (2016).

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