Roles of 3,3′,4′,5-tetrachlorosalicylanilide in regulating extracellular electron transfer of *Shewanella oneidensis* MR-1

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Microbial extracellular electron transfer (EET) is critically involved in many pollutant conversion processes in both natural environment and engineered bioelectrochemical systems (BES), but typically with limited efficiency and poor controllability. In this study, we discover an important role of uncouplers in affecting the microbial energy metabolism and EET. Dose of lower-concentration 3,3′,4′,5-tetrachlorosalicylanilide (TCS) in the anolyte promoted the current generation and substrate degradation of an MFC inoculated with *Shewanella oneidensis* MR-1. However, higher TCS dosage caused obvious microbial inhibition. Our results suggest a previously unknown role of uncouplers in regulating the microbial EET. In addition, the underlying mechanisms of such processes are investigated. This work broadens our view about the EET behaviors of microorganisms in real water environment where uncouplers are usually present, and suggests a possible new approach to regulate microbial EET in BES.

Microorganisms can gain energy from the metabolism-associated electron transfer processes for proliferation and maintenance1–3. In particular, some microbial species have developed the ability to respire external electron acceptors and exert non-negligible influences on processes from geochemical metal cycling to water environmental bioremediation and energy production4–7. Thus, extracellular electron transfer (EET), as a critical foundation of all these processes, needs more attention and careful manipulation2,8. However, such processes in real environment scenario are still poorly understood9,10. Especially, given the usually low EET efficiency and controllability in actual systems11, effective approaches for regulating the microbial EET process are needed.

In metabolism, microbes invest a considerable fraction of generated energy to sustain cell growth and metabolic activity. Under aerobic conditions, the substrate oxidation and electron transfer create a proton motive force (PMF) across the intracellular cytoplasm membrane, which drives the ATP formation via phosphorylation of ADP. Thus, the energy metabolism and electron transport processes are normally coupled by the conservation of free energy in the form of ATP12. However, this coupling can be disrupted or even completely broken by uncouplers to short-circuit the PMF12–14. As a result, the growth of aerobic cells would decrease in the presence of uncoupler due to restrained ATP production and reduced energy for biomass synthesis15–17. However, can the energy metabolism and EET of bacteria under anaerobic conditions also be affected by uncoupler? Even more, could this be adapted as a possible way for microbial EET manipulation? These questions remain unsolved so far.

Herein, we conducted a first investigation into this process by using *Shewanella oneidensis* MR-1 as a model bacterium and 3,3′,4′,5-tetrachlorosalicylanilide (TCS) as a metabolic uncoupler. TCS has been widely adopted as an environmentally-benign uncoupler to reduce yield of activated sludge18,19. Our results show that TCS could also significantly affect the anaerobic metabolism and electron transfer processes. In light of the ubiquity of uncouplers in natural environment, our work implies that the EET processes in natural environments and various engineered systems might need to be reexamined. Moreover, this work might also open up a useful new approach to engineer microbial EET and bioelectrochemical processes.
Results
Electricity generation and EET performances of MFCs. MFC tests were conducted to investigate the effects of TCS on the EET of *S. oneidensis* MR-1. A moderate TCS concentration of 50 μg/L was adopted. As shown in Fig. 1, while no voltage was detected in all the abiotic and dead-cell controls, considerable power generation was observed in all the inoculated MFC, suggesting that the electron generation relies critically on the activities of the inoculated *Shewanella* cells. Notably, the TCS effects on electricity generation were concentration dependent. Addition of 50 μg/L was found to significantly increase the output voltage compared to the TCS-free control, while 400 μg/L led to obvious suppression.

This concentration-dependent impacts of TCS could be more clearly seen from the total recovered charge (Fig. 2). TCS of 50 μg/L lead to the highest quantity of recovered electricity in the MFC, which was over two-fold higher than the TCS-free control. However, a distinct drop of the electricity generation was observed when further increasing the TCS dosage. The recovered electricity at 400 μg/L TCS was even below the control. These results suggest that lower concentration of TCS could promote the microbial EET while a higher concentration may lead to inhibition.

Lactate degradation. The lactate consumption rates, which indirectly reflect the overall microbial activity, also showed a similar trend in response to TCS addition, with the highest removal obtained at 50 μg/L TCS while the lowest achieved at 400 μg/L (Fig. 3). The statistical analysis of the three sets of data showed F and P values of 8.67963 and 0.01695 respectively, suggesting that the lactate consumption rates under the tested conditions are significantly different (p<0.05). Notably, no significant differences in the biomass growth were observed regardless of the different TCS strength (data not shown). Thus, the promotion of lactate consumption at lower TCS dosage was likely associated with a shift of substrate metabolic pathway, as detailed below.

Notably, similar concentration-dependent impacts of TCS (i.e., promoting EET at lower concentrations but inhibiting at higher concentrations) were also observed for several other EABs (data not shown), indicating that TCS could serve as an effective regulator to tune the microbial EET for a wide spectrum of bacteria.

Discussion
The above results demonstrate a concentration-dependent effect of TCS on microbial EET and metabolism. The underlying mechanisms of EET regulation by uncouplers must be different from that by electron mediator, as the latter solely shuttle electrons between bacteria and extracellular electron acceptors20–22.

Fig. 4a depicts the dominant pathways of electron transfer and energy metabolism within *Shewanella* cells under normal conditions (without uncoupler addition). Firstly, the electrons derived from substrate oxidation are transferred to electron-transporting chain (ETC) proteins, from which a part of the electrons are further delivered outside. The ETC also acts as a proton pump to drive protons out by using the energy carried in electrons. As a consequence, accompanied with the electron transfer process, a PMF is built up across the membrane. Such a PMF plays a critical role in ATP synthesis. A part of the protons further diffuse outside the cells, while another part are driven back into the cells through the ATP synthases under the PMF, during which process the energy carried by protons are converted to chemical energy in ATP bonds. Therefore, under this circumstance the microbial electron transfer is coupled with the ATP synthesis process. This kind of metabolic pathway, which is also termed as oxidative phosphorylation, presents a highly efficient way for bacteria to obtain and conserve energy.

However, such a coupling was significantly weakened by TCS. Fig. 4b shows the possible patterns of substrate metabolism, electron transport and ATP synthesis in the presence of TCS. Generally, TCS as an uncoupler may affect the energy metabolism and activity of microbial cells by three mechanisms: 1) directly uncoupling the metabolism through dispersing the PMF and hence reducing the ATP synthesis for energy production19,24; 2) inducing more EPS secretion, especially proteins, so as to resist the toxicity of TCS25.
Similar phenomenon was also observed in the presence of high free chlorine\textsuperscript{25}. The increased secretion of extracellular polymeric substances consumes extra energy and further reduces available energy for microbial growth; 3) TCS at higher concentrations may cause microbial inhibition and even trigger cell lysis\textsuperscript{27,28}. These mechanisms may also apply in our system depending on the TCS dosage and environmental conditions. In addition, several other processes might be trigged by TCS, thereby leading to different substrate metabolism and electron transfer behaviors. Firstly, TCS as a proton carrier could introduce a new pathway of proton transfer across the membrane and weaken the PMF, which favors a more efficient pumping of electrons and protons cross the membrane due to a lower electrochemical resistance. This could explain the increased EET and electricity generation of the MFCs with lower TCS concentration. Interestingly, the TCS led to suppressed ATP synthesis via the oxidative phosphorylation route though, no significant decrease in cell growth was observed in our study. Instead, the lactate uptake of \textit{Shewanella oneidensis} MR-1 was slightly stimulated by the 50 \textmu g/L of TCS. This should be associated with a shift of the metabolic pathway for \textit{S. oneidensis} MR-1 under anaerobic conditions. Due to a suppressed oxidative phosphorylation in the presence uncoupler, the cells are forced to metabolize substrate by a fermentative pathway. The latter process also generates small amount of ATP but is independent of oxidative phosphorylation route.\textsuperscript{29} Thus, in order to compensate for the energy deficiency, the cells may aggressively uptake more substrate due to energy hungry effects\textsuperscript{30,31}. This may, to certain degree, compensate for the decreased biomass synthesis caused by metabolic uncoupling. On the other hand, an over high TCS level (400 \textmu g/L in this case) may severely impair the ATP-synthesis and enzyme activity, thereby suppressing the microbial activity for substrate uptake and electricity generation. This TCS strength is unlikely to induce cell lysis in the short-term operation according to pervious studies\textsuperscript{18,25}, but the long-term impacts are yet to be investigated. Moreover, many of the proposed mechanisms here still need to be validated in future studies.

In summary, this study report a new discovery that microbial EET process could be affected by TCS. It expands our knowledge on the EET behaviors of microorganism in real environments whereby uncouplers like TCS are usually present, and opens up a new possibility for manipulating microbial EET and promoting microbial-catalysis associated applications.

**Methods**

**Inocula and cultivation conditions.** The frozen stocks of \textit{Shewanella oneidensis} MR-1 was provided by Kenneth H. Nealson from the University of Southern California and Xiao Xiang from Shanghai Jiao Tong University. Before test, the strain was inoculated from frozen stocks and grown in LB medium at 30\textdegree C until the exponential phase. A sterile sodium lactate minimal salt medium was used as the cultivation medium and was fed into the anodic chamber. The medium contained (in per liter): 3.36 g sodium lactate, 5.85 g NaCl, 11.91 g Hepes, 0.3 g NaOH, 1.498 g NH\textsubscript{4}Cl, 0.097 g KCl, 0.67 g NaH\textsubscript{2}PO\textsubscript{4}•2H\textsubscript{2}O, plus 0.4 ml of a trace mineral stack solution (containing per liter: 1.5 g NTA(C\textsubscript{6}H\textsubscript{9}NO\textsubscript{6}), 30 g MgSO\textsubscript{4}•7H\textsubscript{2}O, 5 g MnSO\textsubscript{4}•H\textsubscript{2}O, 10 g NaCl, 1 g FeSO\textsubscript{4}•7H\textsubscript{2}O, 1 g CaCl\textsubscript{2}•2H\textsubscript{2}O, 1 g CoCl\textsubscript{2}•6H\textsubscript{2}O, 1.3 g ZnCl\textsubscript{2}, 0.1 g CuSO\textsubscript{4}•5H\textsubscript{2}O, 0.1 g AlK(SO\textsubscript{4})\textsubscript{2}•12H\textsubscript{2}O, 0.1 g H\textsubscript{3}BO\textsubscript{3}, 0.25 g Na\textsubscript{2}MoO\textsubscript{4}•2H\textsubscript{2}O, 0.25 g NiCl\textsubscript{2}•6H\textsubscript{2}O, 0.25 g Na\textsubscript{2}WO\textsubscript{4}•2H\textsubscript{2}O). After autoclaving, the sterilized medium was added with 0.4 ml of filter-sterilized amino acid solution (containing per liter: 2 g L-glutamic acid, 2 g L-arginine, 2 g DL-serine) and 0.4 ml of filter-sterilized vitamin solution (containing per liter: 2.0 g biotin, 2.0 g folic acid, 10.0 g pyridoxine HCL, 5.0 g riboflavin, 5.0 g thiamine, 5.0 g nicotinic acid, 0.1 g cyanocobalamin, 5.0 g p-aminobenzoic acid, 5.0 g thiotic acid).

TCS, a protonophore widely present in the environment and commonly used proton shuflter, was selected as a model uncoupler for the tests. All chemicals used in this study were of analytical grade, and the solutions were prepared with ultrapure water (Millipore Co., USA).

**MFC experiments.** Dual-chamber MFCs with a 110-mL anodic chamber were used to validate the electricity generation capability of the tested bacteria\textsuperscript{26}. Carbon felt (Beijing Sanye Carbon Co., China) with a projected surface area of 8 cm\textsuperscript{2} was used as both the anode and cathode materials. The electrodes were connected via an external...

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**Figure 4** | Pathways of cross-membrane electron transfer, proton transport, and energy metabolism of electrochemical active microorganisms (a) under normal conditions; and (b) with TCS addition. The orange and purple arrows represent the move directions of protons and electrons respectively.
circuit with a 1000-Ω resistance. A proton exchange membrane (GEFC-10N, GEFC Co., China) was used as the separator. The catholyte consisted of 50 mM potassium ferricyanide in phosphate buffer solution (50 mM, pH 7.0). Prior to the experiment, both the anode and cathode chambers were flushed with ultrapure nitrogen gas to ensure an anaerobic atmosphere. The anode chamber was inoculated with S. oneidensis MR-1. After adding 100 ml of sterile mineral salt medium containing sodium lactate as carbon source, a pre-determined concentration of TCS was dosed into the anode chambers of MFCs. Three other MFCs without inoculation were used as the abiotic control, while another three MFCs inoculated with dead cells were used as the dead-cells control. The MFCs were operated in a batch mode at 25 ± 1°C, and all tests were conducted in triplicate.

The voltage across the external resistance was continuously recorded by using a data acquisition switch unit (34970A, Agilent Inc., USA) connected to a computer.

Analysis. Lactate concentration was measured using a high performance liquid chromatograph (HPLC, 1101, Agilent Inc., USA) equipped with an ultraviolet detector and a C18 reversed phase chromatographic column. The mobile phase consisted of 97.5% H2O:PB solution (1 mL 99.5% HPLC-grade H2O:PB in 1 L deionization water, pH 3.0) and 2.5% acetonitrile solution (HPLC grade).

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Author contributions
Y.P.W., S.S.Y. and H.Q.Y. designed the experiments; Y.P.W., S.S.Y., H.L.Z., W.W.L. and Y.Y.C. conducted the experiments; Y.P.W., W.W.L. and H.Q.Y. contributed to the planning and coordination of the project and writing of the paper; All authors contributed to discussion about the results and the manuscript.

Additional information
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