Enhanced antioxidant activity, antibacterial activity and hypoglycemic effect of luteolin by complexation with manganese(II) and its inhibition kinetics on xanthine oxidase

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The present study aims to improve the biological activities of luteolin by complexation with manganese(II). UV-visible spectroscopy, infrared spectroscopy, thermogravimetric analysis and elemental analysis were adopted to assess the relevant interaction of luteolin and manganese(II) ions and the chelation sites. The antioxidant activity, hypoglycemic effect and antimicrobial activity of luteolin–manganese(II) complex with respect to its parent luteolin and the inhibition effect of which on xanthine oxidase were investigated and compared. The spectroscopic data indicated that luteolin reacts with manganese(II) cations through the chelation sites of 5-hydroxy and 4-carbonyl in two luteolin molecules. Antioxidant and antibacterial activity were enhanced after the complexation of manganese(II) cations with luteolin. An inhibition effect assay found that luteolin and luteolin–manganese(II) complex reversibly inhibited xanthine oxidase in a competitive manner. Luteolin–manganese(II) complex had a more remarkable hypoglycemic effect than luteolin by increasing the glucose consumption in liver tissue.

1. Introduction

Luteolin, which is present as glycosylated forms in various kinds of vegetables and fruits, belongs to the flavone subclass of flavonoids.1–3 It has been reported to have anti-inflammatory, antioxidative, antidiabetic, antimicrobial and anticancer activities in vivo.4–8 Luteolin has received increasing attention due to its various biological activities and pharmacological effects and has been widely applied in the fields of medicine and functional foods.9–11 Free radical scavenging and metal chelating are the most common reactions attributed to the antioxidant mechanism.12 Hence, antioxidants of low-molecular-weight and metal chelators have received increased attention both as protectors from disease and as therapeutics.13 Most of the flavonoids are potent metal chelators and possess strong antioxidant and free radical scavenging activity.14 Several flavonoids like rutin, quercetin and luteolin have been proven to be good chelating ligands.15,16 They can interact with metal ions via three domains including the dihydroxy group (3′-OH and 4′-OH) in the B ring, the 3-OH or 5-OH and the 4-CO groups in the C ring (Fig. 1).17

Recently, the development of metal-based compounds with flavonoids has been stimulated and the complexation of flavonoids like quercetin, rutin and luteolin with a large number of metal cations such as Cu(II),18 Zn(II),19 V(II),20 Mo(VI),21 Fe(II),22 Fe(III),23 Sn(II)24 and Cr(III)25 has already been reported. In a previous report, copper–quercetin complex was synthesized and confirmed to be much more effective in free radical scavenging activity than the free quercetin.26 The complex formation between luteolin and vanadium(IV) oxide sulphate monohydrate was investigated in another work.27 It was observed that the free radical scavenging activity and ferric ion reducing potential of luteolin was increased after the formation of complex with vanadium oxide cation.

Despite increasing studies of various metal-flavonoids complex and their biological activities,11,15,17,18,21,22 research about the complexation between luteolin and manganese(II) cation has been limited. Therefore, the purpose of the present study was to examine the complex formation between luteolin and manganese(II) cation and confirm the complexation sites in luteolin by means of UV-visible spectroscopy, infrared spectroscopy, thermogravimetric analysis and elemental analysis. Moreover, the variation of antioxidant property, antibacterial activity and hypoglycemic effect of luteolin after chelation with manganese were investigated. In the meantime, the inhibition effects and inhibition kinetics of luteolin and its mental complex on xanthine oxidase were evaluated. Results obtained in this work are expected to provide significant clues to further exploration of luteolin–manganese(II) complex with enhanced biological activities as a more effective xanthine oxidase inhibitor.

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2. Materials and methods

2.1. Materials and chemicals

Hesperidin (pure: 99%) was purchased from Xi'an Realin Biotechnology Co., Ltd. (Xi'an, China). Manganese acetate (Mn(CH3COO)2), 2,2-diphenyl-1-picrylhydrazyl (DPPH), xanthine and xanthine oxidase were purchased from Sigma Aldrich Chemical Co. (St. Louis, MO, USA). *Escherichia coli*, *Staphylococcus aureus*, *Listeria monocytogenes* and *Pseudomonas aeruginosa* were provided by Guangdong Culture Collection Center (Guangzhou, China). Luria–Bertani (LB) bouillon culture-medium and LB agar culture-medium were purchased from Guangdong Huankai Microbial Science and Technology Co. (Guangzhou, China). Human hepatoma (HepG2) cells (human hepatoma cell line) were purchased from American Type Culture Collection (Rockville, MD, USA). RPMI 1640 medium, Hanks’ Balanced Salt Solution (HBSS), Dulbecco’s modified eagle medium (DMEM), Fetal bovine serum (FBS), phosphate-buffered saline (PBS, pH 7.4), insulin, penicillin, streptomycin, gentamycin, sterile water and other cell culture reagents were purchased from Gibco Life Technologies Co. (Grand Island, NY, USA). All other chemicals of analytical grade were acquired from Guangzhou Chemical Reagent Factory (Guangzhou, China).

2.2. Synthesis of luteolin

Firstly, 10 g of hesperidin were weighed into a round-bottom flask and absolute ethyl alcohol (300 mL) was added to dissolve it. Then 10 mL H2SO4 was added and the mixture was placed in a water-bath at 60 °C for complete reaction. Afterwards, reaction solution was extracted with ethyl acetate (3 × 300 mL) and the organic phase was discarded. Deionized water and 5% NaCl were orderly used to wash the solution. The hesperetin was obtained from the solution by vacuum drying.

The hesperetin (3.5 g) was dissolved in anhydrous pyridine. Then anhydrous AlCl3 (1.5 g) was added and the mixture was kept in a water-bath at 70 °C for complete reaction. After that, anhydrous pyridine was evaporated and the red residue was added with 10% HCl (100 mL) for hydrolyzation. The hydrolysate was also extracted with ethyl acetate (3 × 300 mL) and the organic phase was discarded. After washing by deionized water, the solution was dried through vacuum drying process and the pale yellow powder (eriodictyol) was obtained.

The eriodictyol (3.5 g) was weighed into a round-bottom flask and dissolved with anhydrous pyridine. The I2 (1.2 g) was added for reaction with it. After complete reaction, the anhydrous pyridine was evaporated and the pale yellow residue was obtained through vacuum filtration. Then 100 mL of absolute ethyl alcohol was employed for dissolution and 0.5 g of activated carbon was added. It was kept at 70 °C for 30 min and the luteolin was acquired by filtration.

2.3. Synthesis of luteolin–manganese(II) complex

The luteolin–manganese(II) complex was synthesized according to a reported work with appropriate modification.17 Briefly, 4.0 g of luteolin was weighed and subsequently dissolved in 100 mL
of absolute ethyl alcohol. A saturated solution of Mn(CH₃COO)₂
was added drop wise to the resulting solution (in the molar ratio
of luteolin : Mn⁺² = 2 : 1) under continuous stirring. The pH of
the solution was adjusted to 4 with 1 mmol L⁻¹ of HCl. After
complete reaction, the solid was precipitated and obtained
through filtration. The solid was washed several times with
absolute ethyl alcohol and dried in desiccators over anhydrous
MgSO₄. The luteolin–manganese(II) complex powder was finally
obtained. The proposed synthetic path of the co-ordination
complex was illustrated in Fig. 1.

2.4. Physical measurements
UV-vis spectra of 0.2 mg mL⁻¹ solutions of the luteolin and the
luteolin–manganese(II) complex were obtained in deionized
water by a UV-vis double beam spectrophotometer (UV-1601,
Shimadzu, Kyoto, Japan) using standard 1.00 cm quartz cells.
IR spectra were recorded with a FT-IR (Vertex 70, Bruker,
Rheinstetten, Germany) spectrometer in the range of 500–
4000 cm⁻¹. Briefly, the luteolin and the luteolin–manganese(II)
complex dried completely were thoroughly mixed with KBr
powder, ground and pressed into a 1 mm pellet, then the
prepared samples were determined by FT-IR spectrometer. The
TG–DSC analysis measurement of the luteolin–manganese(II)
complex under the N₂ atmosphere (flow rate: 10 mL min⁻¹) was
performed by a thermal analyzer (STA 449 F3 Jupiter®, Netzsch,
Bavaria, Germany), between the temperatures of 25–800 °C at
a heating rate of 10 °C min⁻¹.

2.5. Elemental analysis
A scanning electron microscope (Nano SEM430, Nova™, Hills-
boro, Oregon, USA) equipped with an energy dispersive spec-
troscopy (EDS, Oxford, UK) was used to perform the elemental
analysis of the luteolin–manganese(II) complex.

2.6. Measurements of antioxidant activity

2.6.1. Analysis of DPPH radical scavenging activity. The
DPPH radical scavenging activity was measured according to
the method of Dehghan & Khoshkam (2012) with some modi-
fications.¹⁴ Briefly, the luteolin and the luteolin–manganese(II)
complex were firstly dissolved in dimethyl sulfoxide (DMSO) to
obtain different concentrations (0.2–5.0 mg mL⁻¹). Then 1 mL
of sample solution was added to 2 mL of DPPH solution
(1 mmol L⁻¹ in methanol). The reaction mixture was shaken
vigorously and then incubated in the dark at room temperature
for 30 min. Finally, the absorbance of the resulting solution was
measured at 517 nm (Aₒ) by the UV-vis double beam spectro-
photometer (UV-1601, Shimadzu, Kyoto, Japan). As a control,
the absorbance of blank solution of DPPH was also determined
at 517 nm (A₀). The DPPH radical scavenging activity was
calculated according to the following equation:

\[
\text{DPPH radical scavenging activity (\%) } = \left( \frac{A_0 - A_t}{A_0} \right) \times 100\%
\]

2.6.2. Analysis of hydroxyl radical scavenging activity. The
hydroxyl radical scavenging activity of the luteolin and the
luteolin–manganese(II) complex was determined as described in
a previous literature.²³

2.7. Evaluation of antimicrobial activity
Antimicrobial activities of the luteolin and the luteolin–man-
ganese(II) complex were assessed using four kinds of bacteria
including Escherichia coli, Staphylococcus aureus, Listeria mono-
cytogenes and Pseudomonas aeruginosa. These bacteria stored at
−80 °C were inoculated to the LB agar culture-medium and then
incubated in a constant temperature incubator at 37 °C for 18–
24 h. The bacterial suspensions were prepared via the incubated
bacteria and subsequently stored in a refrigerator at 0–4 °C.
Afterwards, 0.2 mL of bacterial suspension for each bacterium
was transferred to a new LB agar culture-medium and the
susceptibility paper immersed in sample solution for 12 h was
posted on the medium. Three replicates were performed for
each bacterium. Penicillin and the sterile water were used as the
positive and negative controls, respectively. All the pre-prepared
media were finally incubated in a constant temperature
incubator at 37 °C for 18–24 h and the inhibiting ring was
measured by a vernier caliper. Each sample was calculated
a minimum of three times, and the mean value was calculated
and reported.

2.8. Measurement of the inhibition rate of luteolin and
luteolin–manganese(II) complex on the xanthine oxidase
(XOD)
It has been reported that xanthine can be catalyzed by xanthine
oxidase, resulting in the production of uric acid which has
a characteristic absorption peak at 290 nm.²⁴ So the enzyme
activity in the present work was measured spectrophotometri-
cally by continuously measuring uric acid formation at wave-
length of 290 nm with xanthine as the substrate.²⁵ Briefly,
will a series of assay solutions were prepared by dissolving 2 mL of
xanthine substrate (fixed concentration: 0.6 mmol L⁻¹) and
0.2 mL of various concentrations of luteolin or luteolin–man-
ganese(II) complex in 2.8 mL of sodium phosphate buffer
(0.2 mol L⁻¹, pH 7.5). These assay solutions were shaken
vigorously and then incubated for 30 min in a 25 °C water-bath.
Then the assay was initiated by adding 0.2 mL of xanthine
oxidase solution (0.5 U mL⁻¹). The absorbance at 290 nm was
measured every 30 s. The enzymatic activity was finally calcu-
lated according to the equation reported by a previous
literature.²⁶

In the reaction system, the concentration of xanthine
substrate was fixed at 0.6 mmol L⁻¹. The effects of different
concentrations of luteolin or luteolin–manganese(II) complex
on the enzymatic activity were determined by changing the
concentrations of XOD. The linear equations were then ob-
tained by plotting the reaction velocities (v) and XOD concen-
trations ([x]) in the enzymatic reaction. A linear equation through
the origin indicates the reversible inhibition, while a straight
line which is parallel with the X axis accounts for the irreversible
inhibition. To describe the competitive inhibition mechanism,
the concentration of XOD was fixed at 10 µg mL⁻¹, and the kinetics of the enzyme in the presence of luteolin or luteolin–manganese[II] complex were investigated using double-reciprocal Lineweaver–Burk plots.²⁴

2.9. Cytotoxicity assay of HepG2 cell

HepG2 cells were cultured in RPMI 1640 medium supplemented with 10% FBS, 100 µg mL⁻¹ streptomycin and 100 units per mL penicillin at 37 °C in a humidified atmosphere cultvator with 5% CO₂. The cytotoxicity of luteolin or luteolin–manganese[II] complex on HepG2 cells were assessed by the methylene blue assay.²⁷ Briefly, HepG2 cells were seeded with a density of 8000 cells per well on a 96-well microplate in 100 µL of culture medium per well and subsequently cultured in an incubator at 37 °C for 24 h. The medium was then removed and the HepG2 cells were washed with 100 µL of sterile PBS. After that, 100 µL of the medium containing different concentrations of luteolin or luteolin–manganese[II] complex were added to each well. After another 24 h of incubation, the medium was removed and the HepG2 cells were stained with 50 µL of methylene blue. Afterwards, the staining solution was discarded and the microplate was washed by using deionized water. Then 100 µL of elution buffer was added to each well and the plates were shaken for 20 min. Finally, the absorbance was measured at 570 nm with a microplate reader. A control group without the cell viability.

2.10. Glucose consumption in HepG2

The glucose consumption in HepG2 was determined according to the method described in a previous work with some modifications.²⁸ Briefly, HepG2 cells were seeded with a density of 5000 cells per well on a 96-well microplate and maintained in the DMEM medium at 37 °C for 12 h. Then they were washed and replaced with DMEM medium which contains high glucose (10 mmol L⁻¹), FCS (5%) and insulin (0.5 µmol L⁻¹). Afterwards, HepG2 cells were incubated at 37 °C for 24 h and then washed with serum-free DMEM high glucose. The cells were treated with luteolin or luteolin–manganese[II] complex with different concentrations (1.6, 3.1, 6.3, 12.5 and 25 mmol L⁻¹). The glucose concentrations in cell culture supernatant of each group were determined with glucose assay kit (Jiancheng, Nanjing, China) following the manufacturer’s instruction. The medium and metformin (25 mmol L⁻¹) were used as the control group and the positive control group, respectively.

2.11. Statistical analysis

Results were expressed as mean ± standard deviation (SD) from three replicates for each assay and the data were analyzed by using SPSS 20.0 software (SPSS Inc., Chicago, IL, USA). The analysis of variance (ANOVA) and Tukey’s test were used to compare the differences among various groups and a value of p < 0.05 was considered to be statistically significant.

3. Results and discussion

3.1. UV-visible spectra analysis

The UV-visible spectra of luteolin [a] and luteolin–manganese[II] complex (b) in deionized water are depicted in Fig. 2. In the UV-visible region of luteolin, two major absorption peaks, like most flavones and flavonols, were observed at 356 and 264 nm for band I and band II, respectively. This result was similar to a previous study.²⁶ It has been demonstrated that band I was annotated for the absorption of cinnamoyl system of luteolin, while band II was considered to be associated with the absorption involving benzoyl system.²⁷ In comparison with absorption spectra of luteolin, a significant change was visualized in that of the luteolin–manganese[II] complex due to the red shift of band I, appeared the absorption peak at 414 nm (band III). The shift of about 58 nm after the addition of manganese solution to luteolin confirms that complex formation takes place between manganese and luteolin. Moreover, the displacement of band II was also observed from the absorption peak of 264 nm (band II) to 268 nm (band IV).

It can be inferred from the chemical structure of luteolin that the 5-hydroxy-4-oxo system (Fig. 1) may be the first site to be involved in the complexation process due to the acidic nature of the 5-OH group and more suitable location of 4-CO group.²⁸ It has been reported that high delocalization of oxygen electrons of 5-OH group could facilitate the π electrons delocalization and a big extended π bond system could be formed by the interaction of manganese with 5-OH group of luteolin via electronic redistribution.²⁹,³¹ Thus, the new ring formation in luteolin–manganese[II] complex was caused by the enhanced

![Fig. 2](image-url)
conjugative effect with the inclusion of 5-OH and 4-CO groups in the C ring (Fig. 1). Hence, the bathochromic shift is highly informative for the coordination position in ligand and results obtained also supported the fact that the newly absorption peaks appeared in luteolin–manganese(II) complex were attributable to the presence of 5-OH and 4-CO groups in A and C rings.

3.2. FT-IR spectra analysis

Fig. 3 depicted FT-IR spectra of luteolin (a) and luteolin–manganese(II) complex (b). The presence of ν(Mn–O) stretching vibration peak at 809 cm⁻¹ in the spectrum of luteolin–manganese(II) complex indicated the formation of metal complex via the complexation process, while luteolin exhibited no such band. The characteristic stretching mode of C=O was shifted from 1655 cm⁻¹ of luteolin to 1635 cm⁻¹ of luteolin–manganese(II) complex due to the interaction of luteolin with manganese. It can be explained by the co-ordination of carbonyl oxygen with metal ion that confirms to be the manganese(II) had bonded to 5-OH group of A ring and 4-CO carbonyl group of C ring. In addition, a decrease in vibrational frequency of ν(C–OH) of luteolin–manganese(II) complex, which was shifted from 1369 cm⁻¹ to 1354 cm⁻¹, was observed probably due to the metal coordination involves the deprotonated ortho-phenolic (OH) group on the A ring of luteolin. The band at 1164 cm⁻¹ in luteolin which attributes to the ν(C–O–C) stretching vibration mode remained unchanged in luteolin–manganese(II) complex, indicating that the ring oxygen atom was excluded from involvement in the metal coordination. The absorption peak at 1655 cm⁻¹ was ascribed to the ring ν(C=C) of luteolin and was shifted to 1585 cm⁻¹ in luteolin–manganese(II) complex. This phenomenon illustrated that a more stable ring was formed due to the increase of conjugate effect after the complex formation. Results discussed above confirmed the complexing reaction of luteolin with manganese(II) and were supportive in the sense that the 5-OH group of A ring and 4-CO carbonyl group of C ring were the metal complexation site.

![Fig. 3](image_url)  
**Fig. 3** The IR-spectra of (a) luteolin and (b) luteolin–manganese(II) complex.

3.3. Thermal study of the luteolin–manganese(II) complex

The TG–DSC curves which depict the characteristic thermal events of luteolin–manganese(II) complex were obtained by a thermal analyzer with heating rate of 10 °C min⁻¹. As shown in Fig. 4, the DSC curve for the complex displayed the endothermic peaks that were completely in agreement with the mass losses observed in the TG curve. The first endothermic peak of luteolin–manganese(II) complex at about 106 °C observed in the DSC curve probably due to the dehydration of the free moisture, in accordance with 7% total mass losses in the TG curve. Another endothermic peak around 310 °C with 20% total mass losses possibly be related to the melting point of the luteolin–manganese(II) complex or the dehydration of crystal water or coordinated water. The final thermal decomposition of the luteolin–manganese(II) complex was appeared at above 500 °C with the final mass losses of 82%.

3.4. Elemental analysis of the luteolin–manganese(II) complex

Elemental analysis results of the luteolin–manganese(II) complex are presented in Table 1 and Fig. 5. As shown in Fig. 5, elements of C, O and Mn were observed in the luteolin–manganese(II) complex, confirming the coordination reaction between luteolin and the manganese ions. The weight percentages of C, O and Mn in luteolin–manganese(II) complex were 62.48%, 29.19% and 8.33%, respectively, while the corresponding theoretical values were 59.13%, 31.63% and 9.06%, respectively (Table 1). Moreover, the atomic number percentages were 68.24%, 28.81% and 2.95% respectively, which were also basically consistent with the theoretical values presented in Table 2.

3.5. Antioxidant activities

3.5.1. DPPH radical scavenging activity. The DPPH radical is a stable nitrogen-centered free radical which has been widely adopted for evaluating the free radical scavenging activities of naturally antioxidative compounds and antioxidants. The
DPPH radical scavenging activities of luteolin and luteolin–manganese(II) complex with different concentrations were monitored and the results are shown in Fig. 6A. Significant increases ($p < 0.05$) of the DPPH radical scavenging activity for both luteolin and luteolin–manganese(II) complex were observed with their concentrations in the range of 0–0.8 mmol L$^{-1}$. However, the increase of DPPH radical scavenging activity was not significantly ($p > 0.05$) different in the concentrations of 0.8–1.0 mmol L$^{-1}$. The DPPH radical scavenging activities of 1 mmol L$^{-1}$ luteolin and 1 mmol L$^{-1}$ luteolin–manganese(II) complex reached to 75.24% and 89.14%, respectively. Results confirmed that luteolin was a bioflavonoid with good radical scavenging activity and antioxidant activities. From Fig. 6A, the DPPH radical scavenging activity of luteolin–manganese(II) complex was higher than that of luteolin in the same concentrations. In the DPPH radical scavenging reaction, antioxidants donate the hydrogen or electron to DPPH$^-$ and convert it into DPPH$^-$H, thus the antioxidant activity of compounds mainly depends on the electron- or hydrogen-donating ability. The formation of manganese–luteolin complex electrons increased its scavenging abilities of oxidants and free radicals. Our findings were also in agreement with a previous work that luteolin–vanadium(II) complex showed a better DPPH radical scavenging ability compared to luteolin.

Table 1 Analysis of relative elements mass fraction and relative atoms number of the luteolin–manganese(II) complex

| Elements | Wt$^a$ (%) | Wt (theoretical value, %) | At$^b$ (%) | At (theoretical value, %) |
|----------|------------|---------------------------|------------|---------------------------|
| C        | 62.48      | 59.31                     | 68.24      | 69.77                     |
| O        | 29.19      | 31.63                     | 28.81      | 27.90                     |
| Mn       | 8.33       | 9.06                      | 2.95       | 2.33                      |

$^a$ Wt: the weight percentage. $^b$ At: the atomic number percentage.

Table 2 The diameter of inhibition zone of the luteolin and the luteolin–manganese(II) complex

| Bacteria types              | Sterile water | Luteolin | Luteolin–manganese(II) complex | Penicillin |
|-----------------------------|---------------|----------|--------------------------------|------------|
| Escherichia coli            | 6.02 ± 0.08$^a$ | 7.36 ± 0.12$^b$ | 10.63 ± 0.18$^c$ | 24.35 ± 0.16$^d$ |
| Staphylococcus aureus       | 5.75 ± 0.09$^a$ | 9.56 ± 0.11$^b$ | 12.45 ± 0.16$^c$ | 21.98 ± 0.21$^d$ |
| Listeria monocytogenes      | 5.48 ± 0.06$^a$ | 6.78 ± 0.16$^b$ | 9.49 ± 0.15$^c$ | 12.56 ± 0.12$^d$ |
| Pseudomonas aeruginosa      | 6.12 ± 0.08$^a$ | 6.89 ± 0.14$^b$ | 8.78 ± 0.12$^c$ | 18.98 ± 0.17$^d$ |

$^a$ Data are expressed as mean ± SD with triplicates. Values within the same row with different letters are significantly different at $p < 0.05$. 

3.5.2. Hydroxyl radical scavenging activity. Several diseases such as diabetes is principally induced by hydroxyl radical which has been confirmed to be the most reactive and poisonous free radical. In order to further verify the antioxidant activity of the prepared complex and its parent compound, luteolin and luteolin–manganese(II) complex of different concentrations were also investigated to scavenge the hydroxyl radical generated from sodium salicylate by the hydroxyl radical scavenging assay. As shown in Fig. 6B, the hydroxyl radical scavenging activities increased significantly ($p < 0.05$) to 74.43% and 80.34% respectively after treatment with 1 mmol L$^{-1}$ luteolin and luteolin–manganese(II) complex. In addition, like the DPPH radical scavenging ability, the hydroxyl radical scavenging activity of luteolin–manganese(II) complex was higher than that of luteolin in the same concentrations, indicating the enhanced antioxidant activity of metal–luteolin complex compared to the parent antioxidant, luteolin. It has been reported that the antioxidant activity of flavonoids mainly depends on their molecular structure, and as for luteolin, it is possibly attributed to the prominent contribution of the 5-OH in A ring. The significant change of the chemical properties of the ligand luteolin via the chelation reaction with manganese(II) ions is considered to be the main reason for the enhanced antioxidant activity of the complex.

3.6. Antibacterial activity

Table 2 describes the antibacterial activities of the luteolin and the luteolin–manganese(II) complex. The diameter of inhibiting zone on the LB agar culture-medium was adopted to evaluate the antibacterial activity. As shown in Table 2, all four bacterial strains treated with luteolin and luteolin–manganese(II) complex presented bigger ($p < 0.05$) diameter of inhibiting zone than the negative control, which were supportive for the obvious antibacterial activity of luteolin and luteolin–manganese(II) complex. Moreover, the bigger ($p < 0.05$) inhibiting rings of luteolin–manganese(II) complex on the inoculated mediums in comparison with those of luteolin verified the enhanced antibacterial activity after the complexation of metal ions with luteolin.

3.7. Inhibition effects of luteolin and luteolin–manganese(II) complex on XOD

The inhibition effects of luteolin and luteolin–manganese(II) complex on XOD are displayed in Fig. 7. Sharp increases ($p < 0.01$) of inhibitory rates to XOD were observed for both luteolin and luteolin–manganese(II) complex with the increased concentrations from 0 to 60 μg mL$^{-1}$, followed by a gentle increase from 60 to 100 μg mL$^{-1}$. At the same concentration, the inhibitory rates of luteolin–manganese(II) complex on XOD was higher than that of luteolin. Luteolin has been reported to have a significant XOD inhibitory activity in comparison with allopurinol. The results obtained in the present work showed that luteolin–manganese(II) complex has a higher XO inhibitory activity than that of luteolin, resulting from the complexation reaction of manganese ions with luteolin.

3.8. Determining the inhibition type by Lineweaver–Burk plot analysis

In order to corroborate the reversibility of inhibition, the concentration of xanthine substrate was fixed at 0.6 mmol L$^{-1}$ and plots of the enzymatic activity ($v$) vs. concentration of XOD were constructed as shown in Fig. 8A and B for luteolin (0, 20, 40, 60 and 80 μg mL$^{-1}$) and luteolin–manganese(II) complex (0, 20, 40, 60 and 80 μg mL$^{-1}$), respectively. As depicted in Fig. 8A and B, straight lines for each concentration of luteolin and luteolin–manganese(II) complex were obtained and all passed through the origin. Moreover, the slope of the line was decreased along with the increased concentrations of both luteolin and its complex. Straight lines passing through the origin and decreased slope of the line with increased inhibitor concentrations have been reported to be the obvious phenomenon indicating the reversible inhibition. Therefore, the inhibition of xanthine oxidase by luteolin and its complex was reversible, which was consistent with the findings investigated in a previous literature.

The kinetics of the xanthine oxidase (10 μg mL$^{-1}$) in the presence of luteolin and luteolin–manganese(II) complex were investigated by using double-reciprocal Lineweaver–Burk plots (Fig. 8C and D). It could be clearly observed from Fig. 8C and D that the intercepts of Y axis [1/v] remained unchanged and the intercepts of X axis [1/S] increased along with increasing concentrations of luteolin and luteolin–manganese(II) complex. These results indicated that both luteolin and luteolin–manganese(II) complex induced competitive inhibition, which were in agreement with the findings reported in previous work. This is possibly because that luteolin and luteolin–manganese(II) complex can connected directly with the active position of XOD and compete with the xanthine substrate during the catalytic process.

3.9. Cytotoxicity and glucose consumption assay in HepG2 cell line

Normal amounts of insulin are inadequate to trigger normal insulin response in liver, skeletal muscle and adipose tissue, also known as insulin resistance, is the principle characteristic
of diabetes mellitus.\textsuperscript{40,41} It has been proven that skeletal muscle was the main site of insulin-stimulated glucose uptake \textit{in vivo} and impaired skeletal muscle glucose was the dominating reason leading to the increase of blood glucose in the pathogenesis of insulin resistance and type two diabetes, as glucose transport is the rate-limiting step in muscle glucose metabolism.\textsuperscript{28,42,43} However, liver is the primary organ and liver cells become resistant first for metabolism of glucose, resulting in impaired glycogen synthesis and failure to suppress the production of glucose.\textsuperscript{44} Hence, HepG2 cell lines were employed in the present work to investigate glucose uptake by treatment with luteolin and luteolin–manganese(II) complex, and to elucidate the hypoglycemic effect of them. As shown in Fig. 9A, luteolin and luteolin–manganese(II) had no significant influence on the cell viability of HepG2 in the concentration range of 3.1–25 µg mL\(^{-1}\). Several metal–flavonoid complexes were investigated previously\textsuperscript{31,45–47} and some complexes including the vanadium(II)–flavonoid and oxidovanadium(IV)–flavonoid complexes have been proven to have considerable cytotoxicity on cancer cells.\textsuperscript{17,22,46,47}

![Fig. 8](image1.png)  
**Fig. 8** Plots of enzyme activity (v) vs. concentration of xanthine oxidase (S) for different concentrations of luteolin (A) and luteolin–manganese(II) complex (B). The concentration of xanthine substrate was 0.6 mmol L\(^{-1}\). Lineweaver–Burk plots of (1/v) vs. (1/S) for different concentrations of luteolin (C) and luteolin–manganese(II) complex (D). The concentration of xanthine oxidase was 10 µg mL\(^{-1}\).

![Fig. 9](image2.png)  
**Fig. 9** Effects of luteolin and luteolin–manganese(II) complex on cell viability (A) and glucose consumption (B) of HepG2 cells. Values are presented as means ± SD with triplicates. * \(p < 0.05\), ** \(p < 0.01\) and *** \(p < 0.001\) respectively compared with the control, \(+p < 0.05\), \(;++p < 0.01\) and \(;+++p < 0.001\) by Tukey's test.
After treated with different concentrations of luteolin and luteolin–manganese(II) complex, the glucose consumption by HepG2 cells increased remarkably compared with the control. In particular, the glucose consumption showed a highly significant increase ($p < 0.001$) with the concentrations higher than 6.3 μg mL$^{-1}$. Furthermore, in comparison with luteolin, glucose uptake by HepG2 cells treated with luteolin–manganese(II) complex were significant increased at the same concentrations of 1.6, 3.1 and 6.3 μg mL$^{-1}$ ($p < 0.05$), 12.5 μg mL$^{-1}$ ($p < 0.01$) and 25 μg mL$^{-1}$ ($p < 0.001$), separately. Results of this test showed that hypoglycemic effects of luteolin and luteolin–manganese(II) complex were obviously achieved by increasing the glucose consumption in liver tissue and luteolin–manganese(II) complex had a more remarkable hypoglycemic effect.

4. Conclusions

The metal complex of manganese(II) with luteolin was synthesized by means of the UV-visible spectroscopy, infrared spectroscopy, thermogravimetric analysis and elemental analysis. Results assumed that the –CO group of the ligand was coordinated with one of the neighboring –OH group in luteolin. Luteolin was suggested to react with manganese(II) cation through the chelation sites of 5-hydroxy and 4-carbonyl in two luteolin molecules from the spectroscopic data. The luteolin–manganese(II) complex was confirmed to have higher biological activities including antioxidant activity, antibacterial activity and hypoglycemic activity in comparison with the parent luteolin. Luteolin and luteolin–manganese(II) complex reversibly inhibited xanthine oxidase in a competitive manner and the chelation of manganese(II) ions by luteolin showed stronger inhibitory activity on xanthine oxidase than nature luteolin.

Conflicts of interest

The authors declare that they have no conflicts of interest.

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