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Abstract:

22 A colorimetric immunoassay based on blue silica nanoparticles (Blue-SiNPs) was developed 23 for quantitative detection of *Salmonella pullorum* (*S. pullorum*)*.* In this method, Blue-SiNPs were 24 synthesized by doping C.I. reactive blue 21 into silica nanoparticles using a reverse 25 microemulsion method. Blue-SiNPs functionalized with the anti-*S. pullorum* were employed as 26 detector probe. Magnetic nanoparticles (MNPs), employed as supports for the immobilization of 27 polyclonal antibody against *S. pullorum*, were used as capture probe. The sandwich structures of 28 MNPs-*S. pullorum*-Blue-SiNPs were separated by magnet and etched by NaOH. The C.I. reactive 29 blue 21, released from silica nanoparticles, was used as a colorimetric indicator. The absorbance 30 of C.I. reactive blue 21 at 675 nm is proportional to the concentration of *S. pullorum*. Under the 31 optimal conditions, the developed colorimetric immunoassay exhibited a wide dynamic range of 4.4×10^2 CFU/mL to 4.4×10^7 CFU/mL toward *S. pullorum* with a detection limit of 4.4×10^1 33 CFU/mL. For application of the assay, this method is not influenced by complex matrix of 34 practical samples, The recoveries of *S. pullorum* from chicken liver samples were from 94.5% to 35 108% with a good correlation coefficient (R^2 =0.9989) with those obtained by an official standard 36 culture method. We also show that this colorimetric immunoassay can be carried out on a 37 microplate reader with a 96-well plate. The method is particularly economic, simple, rapid, 38 specific and good stability. The new technology provided a basis for the detection of other 39 pathogenic bacteria and viruses. Such a simple colorimetric immunoassay holds great potential as 40 on-site tool for clinical diagnosis of bacteria and viruses.

Introduction

45 Conventional bacterial testing methods involve several basic steps: pre-enrichment, selective 46 enrichment, selective plating, and biochemical screening and serological confirmation¹. Culture 47 medium preparation, inoculation of plates and colony counting makes these methods labor 48 intensive and time consuming, it may take several days before the results are obtained². There is 49 increasing demands for the rapid and reliable detection of food borne pathogens. Advent of 50 biotechnology has greatly altered pathogenic bacteria testing methods. Colorimetric immunoassay 51 is one of the widely used among all assay techniques, taking advantage of antigen-antibody 52 reaction and color change to identify a substance³. It has gained great attention in the various 53 research areas including food safety, clinical diagnosis and environmental monitoring^{4, 5}. In 54 comparison with fluorescence-based assay, colorimetric immunoassay has significant advantages 55 of low cost without the requirement of fluorescent-labeled antibodies and expensive instruments⁶. 56 Colorimetric immunoassay can be used for on-site detection with a portable UV-Vis absorption 57 spectrometer.

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58 Transforming the detection event into color change is a key technology of colorimetric 59 immunoassays, which is very crucial for obtaining low detection $\lim_{t \to 9} t^{7-9}$. Enzyme-labeled 60 nanoparticles and gold nanoparticles are commonly exploited as signal transduction tools in 61 colorimetric immunoassays^{10, 11}. The carried enzyme molecule is entered and participated in the 62 catalytic reaction, and the high efficiency of enzymes makes them especially suitable for 63 ultrasensitive bioanalysis. However, short lifetime, high price and the critical operating situation 64 limit enzyme immunoassay applicability¹². In recent years, the great superiority of gold 65 nanoparticles in the colorimetric immunoassays has been reported. Gold nanoparticles have been 66 used in colorimetric immunoassays to enhance the stability, improve sensitivity, or simplify the 67 manipulation process. The color and absorption spectrum of gold nanoparticles shift to longer 68 wavelengths if aggregation occurs. However, the absorption intensity of gold nanoparticles is 69 substantially affected by various factors such as pH, temperature, and salt concentration^{13, 14}. 70 Furthermore, it is not economical to use gold nanoparticles for the routine detection of bacteria in 71 foods. As the reported assay is imperfect, we aim to develop a colorimetric immunoassay for

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72 pathogenic bacteria detection, which have the potential to fulfill the WHO ASSURED criteria 73 (affordable, sensitive, specific, user friendly, robust and rapid, equipment free, deliverable to those 74 who need them).

75 In addition to the above-mentioned nanoparticles, silica nanoparticles (SiNPs) are widely 76 used in several fields including disease labeling, drug delivery and biosensor¹⁵. But most reported 77 SiNPs are white or colorless, and they are not suitable for signal conversion and signal 78 amplification¹⁶. Organic dye has rich color and good stability, and does not fade in the harsh 79 conditions of light, heat, acid, alkali and so on. From a chemical point of view, so many kinds of 80 organic dyes and their good chemical reactivity makes them very suitable for dyeing silica 81 nanoparticles to synthesis colored silica nanoparticles. Colored silica nanoparticles are a good 82 candidate optical label in the biotechnological systems due to their inherent advantages, such as 83 bright color, easy preparation and good biocompatibility^{17, 18}. In our previous work, we had 84 developed an agglutination test for simultaneous detection of two different pathogenic bacteria 85 using the colored silica nanoparticles as carrier .

86 Practical samples are highly complex consisting of fats, proteins, minerals and even 87 sometimes contain antimicrobial preservatives. In order to discriminate the target pathogen from 88 complex matrix, a separation step is normally required. Further, they are numerically very low, so 89 efficient pathogen separation and concentration techniques need to be evolved for specific 90 detection of pathogens and to avoid false-negative results²⁰. Magnetic nanomatarials, due to their 91 unique properties including being conveniently manipulated by a magnet, high surface to volume 92 ratio, and fast kinetics in solution, have been widely applied to rapid, efficient, and specific 93 capture and enrichment of target bacteria from the original samples^{21, 22}. For example, Pang's 94 group had successfully used antibody-modified magnetic nanomaterials to capture *Salmonella f typhimurium* at ultralow concentration (∼10 CFU/mL)²³. On the other hand, magnetic capture can 96 easily be coupled with many detection methods, such as colorimetric immunoassay, PCR, 97 fluorescence observation, electrochemical immunosensor, When MNPs are applied to colorimetric 98 immunoassays, the assay is usually fast and simple to operate, which offers a promising platform 99 . for the detection of various analytes 24 .

100 The aim of this work is to exploit blue silica nanoparticles (Blue-SiNPs) and magnetic

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101 nanoparticles (MNPs) based colorimetric immunoassay for the rapid detection of pathogenic 102 bacteria. In this study, *Salmonella pullorum* (*S. pullorum*) was used as a model analyte. In most 103 developed nations, *S. pullorum* is a common infectious pathogen and can result in acute systemic 104 disease and a high incidence of mortality in young poultry ²⁵. Rapid and sensitive detection of *S*. *pullorum* is of great importance for poultry breeding. Aimed to make a stable color indicator for 106 sandwich immunoassay, we chose organic dye (C.I. Reactive Blue 21) to synthesize the Blue-107 SiNPs using an inverse microemulsion method. Organic dyes were modified to the surface of 108 SiNPs by silane coupling agent. Herein, magnetic nanoparticles (MNPs) and Blue-SiNPs were 109 respectively modified with antibodies to construct antibody-coated magnetic nanoparticles (IgG-110 MNPs) and antibody-coated Blue-SiNPs (IgG-Blue-SiNPs). The IgG-MNPs were used for 111 enrichment and separation of *S. pullorum*. The IgG-Blue-SiNPs, which encapsulated hundreds of 112 C.I. Reactive Blue 21 into a single nanosphere, were employed as reporter probes to provide a 113 highly amplified and stable signal. After adding sodium hydroxide in the detection solution, Blue-114 SiNPs generate a colorimetric signal that is directly proportional to the concentration of *S. pullorum*.

Experimental Section

Materials and Instruments

118 A variety of bacteria were employed in this work including *S. pullorum* as the target bacteria, 119 and *Bacillus subtilis* (*B. Subtilis*), *Escherichia coli* (*E. coli*), *Staphylococcus aureus (S. aureus*), *Enterobacter sakazakii* (*E. Sakazakii*) as control group. All bacteria were conserved by the 121 laboratory of author. *S. pullorum* was grown at 37 °C in Lysogeny broth (LB) medium with 122 shaking. Cells were harvested in late exponential growth phase by centrifugation and washed in 123 triplicate using physiological saline aqueous solution. Concentration of the bacteria was confirmed 124 by the colony counting (CFU/mL). The enriched bacterial were inactivated with 0.5% 125 formaldehyde and stored at 4 °C before use.

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126 Triton X-100, cyclohexane, hexanol, glutaraldehyde (GLU, 50 wt %) and ammonia (25-28 127 wt%) were obtained from Chengdu Kelong Chemical Reagent Co., Ltd. (Chengdu, China). 3-[2- 128 (2-Aminoethylamino) ethylamino] propyl-Trimethoxysilane (APTMS), Tetraethyl orthosilicate 129 (TEOS) were obtained from Aladdin Industrial Inc. (Shanghai, China). Bovine serum albumin

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130 (BSA) was obtained from Beijing Dingguo Biotechnolgy Co. Ltd. (Beijing, China). Magnetic 131 nanoparticles (Prot Elut NHS, 100-120nm, MNPs,) were obtained from Enriching Biotechnology 132 Ltd. (Shanghai, China). C.I. Reactive Blue 21 was supplied by Zhejiang Shunlong Chemical Co., 133 Ltd (Zhejiang, China). Polyclonal antibody against *S. pullorum* (100µg/mL) was purchased from 134 China Institute of Veterinary Drugs Control (Beijing, China). Other reagents were all of analytical 135 grade and were used as received without further purification. The water used was doubly distilled. 136 Hitachi SU-70 scanning electron microscopy (SEM) was purchased from Hitachi Inc. (Tokyo, 137 Japan). Malvern Nano 2S potential laser particle analyzer was provided by Malvern Instruments 138 Co., Ltd. (Worcestershire, UK.). Nicolet 380 Fourier transform infrared spectrometer (FTIR) 139 (Thermo, Shanghai, China). SpectraMax 190 microplate reader was purchased from Molecular 140 Devices (Orleans, USA). 3-18K high speed refrigerated centrifuge was purchased from Sigma 141 Laborzentrifugen GmbH (Osterode, Germany). Scientz-09 pat type sterile homogenizer was 142 provided by NingBo Scientz Biotechnology Co.,Ltd (Ningbo, China). All electrochemical 143 experiments were performed at room temperature (25±1 °C).

Synthesis of Blue-SiNPs and Blue-SiNPs-NH²

145 Synthesis of Blue-SiNPs was carried out according to method described by a previous paper 146 with little change²⁶. The details of the procedure are described in the following. The water-in-oil 147 (W/O) microemulsion was prepared at room temperature first by mixing 2 ml surfactant Trition X-148 100, 8 ml oil phase cyclohexane and 2 ml cosurfactant n-hexanol. 0.15 ml C.I. Reactive Blue 21 149 solution (100 mg/mL) and 400 μ L of water were then added. Then the resulting mixtures were 150 homogenized with magnetic force stirring for 15 min to form a W/O microemulsion. In the 151 presence of 100 µL of TEOS and 20 µL APTMS, a hydrolyzation reaction was initiated by adding 152 100 µL of NH3 ·H2O (25-28 wt%) under stirring. The reaction was allowed to stir for 48 h. After 153 the reaction was completed, acetone was added to break the microemulsion and recover the 154 particles. The contents were then centrifuged and washed with ethanol and water several times to 155 remove surfactant molecules and physically adsorbed C.I. Reactive Blue 21 from the particles' 156 surface. The synthesized Blue-SiNPs were characterized by SEM.

157 30 mg Blue-SiNPs were ultrasonically resuspended in the mixed solution of 15 mL ethanol 158 and 40 µL APTMS, it was allowed to complete the silanized reation under stirring overnight at

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159 room temperature. After this reaction, amino groups were introduced to the surface of Blue-SiNPs. 160 Amino-modified Blue-SiNPs (Blue-SiNPs-NH2) were isolated by centrifuging at 8000 rpm and 161 washed three times with 0.01M Phosphate buffered saline solution (PBS, PH7.3). The synthesized 162 Blue-SiNPs and Blue-SiNPs-NH2 were characterized by FTIR, Zeta potential and by SEM.

Surface functionalization of Nanoparticles

164 Covalent immobilization of the antibody onto functionalized Blue-SiNPs surface

165 The polyclonal antibody against *S. pullorum* was directly immobilized onto the functionalized 166 Blue-SiNPs with well-established glutaraldehyde method^{27, 28}. The immobilization protocols were 167 the following (Fig.1): (1) 20 mg of Blue-SiNPs-NH2 was dispersed into 5 mL PBS (0.01M, 168 PH7.3) buffer containing 2.5% glutaraldehyde for about 3h; (2) the nanoparticles were separated 169 by centrifugation and washed with PBS three times. After the nanoparticles re-dispersed in PBS, 170 they were further incubated with anti-*S. pullorum* polyclonal antibody for 2 h at room temperature 171 with shaking; (3) the antibody-coupled Blue-SiNPs(IgG-Blue-SiNPs) were centrifuged at 8500 172 rpm and washed with PBS several times to remove excess antibody and kept at 4 °C in PBS 173 (0.01M, PH7.3) for its following use.

174 Preparation of IgG-MNPs

175 The polyclonal antibody against *S. pullorum* was covalently conjugated to carboxyl-modified 176 MNPs according to the manufacturer's instructions. 100 µL of magnetic nanoparticles (10 mg/mL) 177 were mixed with 50 µL of antibody against *S. pullorum*. The reaction was allowed to proceed at 178 4 °C overnight. The mixture was washed for three times with washing buffer in a magnetic field. 179 Unreacted active groups on the MNPs were blocked with 1% BSA. Finally, the antibody modified 180 magnetic nanoparticle (IgG-MNPs) were dispersed in 1 mL of PBS and stored at 4 °C before use.

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181 Procedures of Colorimetric Immunoassay Measurement

182 The colorimetric immunoassay was carried out as follows: 20 µL IgG-MNPs (1 mg/mL) 183 were mixed with 1 mL *S. pullorum* solution in centrifuge tube. After incubating for 30 min at 184 37°C with gentle shaking, the resulting mixture was separated magnetically and washed with PBS 185 for three times to remove any unbound species. The immune complexes of IgG-MNPs and *S. pullorum* were then dispersed in 20 µL PBS. 20 µL IgG-Blue SiNPs (10 mg/mL) were added into 187 the mixture and subjected to react for 15 min. IgG-Blue SiNPs and IgG-MNPs formed sandwich

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188 structure with *S. pullorum* via immune reaction. The final sandwich immune complexes were 189 washed by PBS to remove unbound IgG-Blue SiNPs. The final sandwich immune complexes were 190 incubated with 50 µL of NaOH aqueous solutions (5 mol/L) for 15–60 min. After the C.I. Reactive 191 Blue 21 was desorbed, the supernatants including C.I. Reactive Blue 21 were separated and 192 transferred to 96-well plate. The absorbance of C.I. Reactive Blue 21 was measured with 193 microplate reader at 675 nm.

Results and discussion

Characterizations of Blue-SiNPs

196 Inverse microemulsion method was chosen to prepare Blue-SiNPs in this study. Inverse 197 microemulsion method is a simple and diverse preparation method for synthesis of silica 198 nanoparticles in laboratory, and it is easy to control the morphology of silica nanoparticles. The 199 physical image of Blue-SiNPs showed that they were bright color and good dispersion in aqueous 200 solution (Fig. 2 inset). The size and morphology of Blue-SiNPs were characterized by SEM (Fig. 201 2). The nanoparticles had excellent monodispersibility and all Blue-SiNPs showed a spherical 202 shape and smooth surface. The average diameters of the nanoparticles determined by SEM were 203 approximately 45±5 nm and the size distribution was also quite uniform and the characteristics of 204 colored-SiNPs were in accordance with descriptions by Tan et al^{29} .

205 The presence of chemical group on the outermost layer of Blue-SiNPs was confirmed by 206 Zeta potential determination. Zeta-potential measurement was carried out using Zetasizer. For 207 determination of Zeta potential, pH of samples was adjusted by addition of 0.01 M HCl or 0.01 M 208 NaOH. All values shown in this work were average of three measurements. Figure 3a displayed 209 Zeta potential of Blue-SiNPs and Blue-SiNPs-NH2 as a function of pH. The isoelectric point (IEP) 210 of Blue-SiNPs is at pH 4.8. When Blue-SiNPs is in the environment of a neutral solution, surface 211 potential of Blue-SiNPs is about -13 mV, which indicates that the Blue-SiNPs surface with a 212 negative charge, because of the presence of hydroxyl groups; The IEP of Blue-SiNPs-NH₂ is 213 shifted to PH 9.5, and in neutral solution, surface potential of Blue-SiNPs-NH₂ is about 18 mV, 214 which indicates that the Blue-SiNPs-NH2 surface with a positive charge. The increase of Zeta 215 potential is attributed to the increasing number of protonated amine groups on the Blue-SiNPs-216 NH₂ surface.

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217 FT-IR analysis provides direct proof for the amine fictionalization. Chemical compositions 218 on the surface of Blue-SiNPs and Blue-SiNPs-NH2 were examined by FTIR (Fig. 3b). Dried 219 samples were measured using KBr pellet method in the range of 400–4000 cm⁻¹. A strong IR 220 absorption bands at the region $980-1220 \text{ cm}^{-1}$, corresponding to the Si–O–Si of the silica core, is 221 found in both Blue-SiNPs and Blue-SiNPs-NH₂. A new band at 2945 cm^{-1} in the Blue-SiNPs-NH₂ 222 is assigned to the N–H of the silica. Compared to Blue-SiNPs, the absorption spectra of Blue- SiNPs-NH₂ has a significantly difference in the region $2900-3450 \text{cm}^{-1}$. These results are 224 consistent with the Blue-SiNPs being successfully coated with an aminenated silica shell. APTMS 225 is thus believed successfully introduced onto the surface of the silica modified nanoparticles.

Principle of Blue-SiNPs-Based Colorimetric Assay for the Detection of *S. pullorum*

227 The principle of Blue-SiNPs-based colorimetric assay for the detection of *S. pullorum* is 228 presented in Fig. 1. We use the colorimetric immunoassay to detect *S. pullorum* based on Blue-229 SiNPs and MNPs. In a typical assay, Blue-SiNPs probes and MMP formed a sandwich structure 230 with the target *S. pullorum* according to the principle of antigen-antibody binding. A magnetic 231 field was used to effectively remove unbound Blue-SiNPs probes. In order to quantify the amount 232 of *S. pullorum*, desorption experiments were carried out. The NaOH aqueous solution (5 mol/L) 233 was used for the release of the loaded C.I. Reactive Blue 21 by etching Blue-SiNPs. There was a 234 correspondence between absorbance of C.I. Reactive Blue 21 and the *S. pullorum* concentration. 235 By measuring the absorbance, we could quantitatively determine the concentration of *S. pullorum* 236 in sample. Meanwhile, we could also qualitatively determine the *S. pullorum* level by the change 237 in visible color. As shown in Fig. 1b, the spectrum of C.I. Reactive Blue 21 exhibited a strong 238 absorption peak at 675 nm. In order to sensitively quantify the amount of *S. pullorum*, we 239 measured the absorbance of C.I. Reactive Blue 21 at 675 nm for establishing a standard curve 240 between the absorbance and the amount of *S. pullorum*.

Optimization of Experimental Conditions

242 To achieve an optimal analytical performance, the experimental conditions for Blue-SiNPs-243 based colorimetric system were further optimized through adjusting incubation times of IgG-244 MNPs with *S. pullorum* from 0 to 60 min, IgG-MNPs concentration from 0.25 to 4 mg/mL. The 245 performance of the developed colorimetric immunoassay could be greatly affected by incubation

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246 times. Fig. 4a shows the absorbance changes of C.I. Reactive Blue 21 with different incubation 247 time, the absorbance rapidly increased with increasing incubation time, which indicates that *S. Pullorum* gradually binds with the magnetic beads via antigen-antibody reaction. And a maximum 249 was attained at 30 min. Further increase incubation time, e.g. 40 min, had very little additional 250 beneficial effect. Besides, the limited diffusion of IgG-MNPs, 30 min incubation was chosen in 251 this study.

252 The concentration of IgG-MNPs in the detection system was optimized by using 20µL IgG-253 MNPs of different concentration from 0.25 to 4 mg/mL. As seen from Fig. 4b, the absorbance 254 increased with the increasing concentration of IgG-MNPs until reach to 1 mg/mL, and the 255 absorbance tended to level off when the concentration is higher than 1 mg/mL. In the condition of 256 1 mg/mL, IgG-MNPs could combine the suitable concentration of *S. pullorum*. Therefore, 1 257 mg/mL was chosen as the optimal concentration of IgG-MNPs for detection of *S. pullorum*in in 258 the following experiment.

Sensitivity of Colorimetric Immunoassay

260 Under optimal experimental conditions, we examined the performance of the proposed immunoassay with different concentrations of *S. pullorum* in the range from 4.4×10^{1} to 4.4×10^{9} 262 CFU/mL. The plot of the absorbance of C.I. Reactive Blue 21versus the logarithm of *S. pullorum* 263 concentration is shown in Fig. 5b. According to the results of absorbance measurements, a linear 264 dependence between the absorbance and the logarithm of *S. pullorum* concentration was obtained 265 in the range from 4.4×10^2 to 4.4×10^7 CFU/mL. The linear regression equation was y = 0.06448 log 266 $x - 0.1052$ ($R^2 = 0.9851$). The *S. pullorum* concentration in the samples was obtained 267 quantitatively via the linear regression equation. The limit of detection (LOD) was 4.4×10^{17} 268 CFU/mL defined as 3SD above the zero-dose response (n= 10). This was comparable with those 269 of other nanolabeled-based immunoassays for pathogenic bacteria. The sensitivity of the classic 270 ELISA reported by Forsythe and Cudjoe was 10⁵ CFU/mL for *Salmonella*³⁰. The higher sensitivity 271 exhibited by the Blue-SiNPs-based immunoassay was attributed to using Blue-SiNPs as a signal 272 amplifier. Each Blue-SiNPs could accumulate thousands of C.I. Reactive Blue 21 molecules 273 owing to its high surface-to-volume ratio. Thus one immunoreaction event could bring multiple 274 C.I. Reactive Blue 21 molecules, leading to large amplification of signals.

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275 The analytical performance of the proposed immunosensor was compared with that of other 276 pathogenic bacteria sandwich immunosensor reported (Table 1). The comparison suggests 277 superior analytical performance of the present immunosensor over some previously reported 278 sandwich immunosensor. The fluoresent immunosensor reported by Pang et al. was an exception²³. They used a fluorescence microscope thus leads to four times of magnitude 280 improvement insensitivity for bacteria detection, the but the technology was more complicated 281 and the time of process that were needed was longer. Magalhaes et al. had also reported a more 282 sensitive method than this work, however the linear range of their method was poor, it could not 283 be used for quantitative detection of *Salmonella typhimurium*.

Specificity of the Blue SiNPs Based Colorimetric Assay

285 The selection of antibody with high specificity is important in developing immunoassay, 286 because the specificity for the measurement of analytes in all immunoassay systems is dependent 287 on the antibody used¹⁰. In this study, a commercial antibody to *S. pullorum* was used. The 288 specificity of the developed colorimetric method was evaluated with five control experiments that 289 involved *B. Subtilis*, *E. coli*, *S. aureus*, *E. Sakazakii* and PBS, respectively. Equal concentrations 290 of *S. pullorum*, *B. Subtilis, E. coli, S. aureus* and *E. Sakazakii* were detected with the same 291 colorimetric immunoassay as mentioned in Section 2.5. After incubation and separation, the 292 colorimetric assay displayed both an obvious blue color (inset in Fig. 6) and a strong increase in 293 absorbance at 675 nm (as showed in Fig.6) in the presence of *S. pullorum*. In contrast, there was 294 neither blue color (inset in Fig. 6) nor strong absorbance (Fig. 6) observed in the presence of *B. Subtilis, E. coli, S. aureus* and *E. Sakazakii* as compared with the PBS without bacteria. These 296 results indicated the high specificity of the proposed method for the detection of *S. pullorum*.

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Applicability of the Assay

298 To evaluate the detection capability and reliability of developed colorimetric immunoassay in 299 practical sample, the commercial chicken livers were used to make artificially contaminated 300 samples. Chicken livers were collected from a local supermarket in Hangzhou (Zhejiang, China), 301 and they were homogenized by a pat type sterile homogenizer. Chicken livers were seeded with 302 serial dilutions of *S. pullorum* to achieve a final concentration from 4.4×10^2 to 4.4×10^7 . The 303 recoveries and CVs of the above-mentioned samples spiked with different concentrations of *S.*

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pullorum are listed in Table 2.With a simple and rapid detection procedure using the colorimetric 305 immunoassay, most of the recoveries of *S. pullorum* for these samples were 101%, 94.5%, 95.0%, 306 104.9%, 102%, 108% respectively, All the recoveries were in the range from 90 to 110%. The 307 correlation coefficient (R^2) of the results obtained from the colorimetric immunoassay was 0.9989. 308 Conventional microbiological methods based on standard plate count are time consuming for 309 identification and quantification of *S. pullorum*. The colorimetric immunoassay established in this 310 study is based on the specific induction of the Blue-SiNPs, resulting in an absorbance change in a 311 dose dependent manner. This assay, which is of high sensitivity and capable of quantitatively 312 recognizing *S. pullorum*, is more sophisticated and specific than the conventional microbiological 313 method. Further, this method is easy to perform and is suitable for detection of *S. pullorum* in 314 practical sample such as animal tissues.

Stability

316 The long-term stability of IgG-Blue-SiNPs and IgG-MNPs was examined. Their stabilities 317 were studied by detecting *S. pullorum* with the developed colorimetric immunoassay after IgG-318 MNPs and IgG-Blue-SiNPs were stored at 4 °C for 1, 7, 30, 60 and 90 days. As shown in Fig.7, 319 during three-month test, the absorbance of the organic dye was almost unchanged for the 90 days' 320 measurement. So these two kinds of immune nanoparticles were found to be able to retain similar 321 reaction activity after storage at 4°C for at least 90 days. Actually, the thus-prepared colorimetric 322 immunoassay is more stable compared with, biosensor based on enzyme-coated nanoparticle³¹. 323 We speculate that the long-term stability mainly attributed to the following two issues: (i) the 324 enzyme labeled antibody was replaced by antibody to modify the Blue-SiNPs, and (ii) anti-*S. pullorum* molecules were covalently immobilized on the surface of the Blue-SiNPs and MNPs.

Conclusions

327 In conclusion, this work demonstrates the development of an advanced colorimetric 328 immunoassay based on IgG-Blue-SiNPs for the detection of *S. pullorum.* Although blue SiNPs 329 was synthesized with inexpensive commercial organic dyes and TEOS, they were excellent 330 markers for colorimetric. Another important innovation present in this paper is the demonstration 331 that the Blue-SiNPs based assays can be carried out in a 96-well plate, thus meeting demands of 332 high-throughput analysis. We show that the direct quantitation of *S. pullorum* by measuring the

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333 absorbance of organic dye is an easy, fast, and reliable way to quantify total bacteria. For 334 application of the assay, this method is not influenced by complex matrix of practical samples and 335 has good sensitivity. Compared with the conventional colorimetric immunoassay system, 336 Highlights of this work mainly are as the follows: IgG-Blue-SiNPs were used for loading anti-*S. pullorum* polyclonal antibody and used as the signal amplification probe, we can directly take 338 advantage of the absorbance of organic dye modified in the surface of SiNPs to quantify the 339 bacterial; this developed colorimetric immunoassay method does not need the catalysis of 340 enzymes to make it more economical and can be applied to more detection places; this method 341 had high selectivity and sensitivity, it could effectively discriminate *S. pullorum* from other four 342 pathogenic bacteria, besides, the detection limit is 4.4×10^{1} CFU/mL, it is much lower than the 343 detection limit of 10^4 CFU/mL of conventional colorimetric immunoassay to detect salmonella. 344 Furthermore, the blue silica nanoparticle and magnetic nanoparticle-based colorimetric 345 immunoassay method does not require sophisticated instruments and is well suitable for other 346 pathogenic microorganisms or even investigated for rapid detection of the virus.

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 $\begin{array}{c} 4 \\ 5 \\ 6 \end{array}$

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437 Fig. 1

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- 439 Fig. 2.
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454 SV Stripping voltammetry, QDs Quantum Dots, AuNPs gold nanoparticles, HRP horseradish

455 peroxidase, IFNS immunofluorescent nanospheres, ABTS 2'-Azinobis-(3-ethylbenzthiazoline-6-

456 sulphonate).

457 Table.1.

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462 Fig.6.

 $\mathbf 1$

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 $\mathbf 1$

467 Table.2.