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# Excess imidacloprid exposure causes the heart tube malformation of chick embryos

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# Abstract

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As a neonicotinoid pesticide, imidacloprid is widely used to control sucking insects on agricultural planting and fleas on domestic animals. However, the extent to which imidacloprid exposure has an influence on cardiogensis in early embryogenesis is still poorly understood. In vertebrates, the heart is the first organ to be formed. In this study to address whether or not imidacloprid exposure affects early heart development, the early chick embryo has been used as an experimental model because of the accessibility of chick embryo at its early developmental stage. The results demonstrate that exposure of the early chick embryo to imidacloprid caused malformation of heart tube. Furthermore, the data reveal that down-regulation of GATA4, Nkx2.5 and BMP4 and up-regulation of Wnt3a led to aberrant cardiomyocyte differentiation. In addition, imidacloprid exposure interfered with basement membrane (BM) breakdown, E-cadherin/Laminin expression and mesoderm formation during the epithelial-mesenchymal transition (EMT) in gastrula chick embryos. Finally, the DiI-labeled cell migration trajectory indicated that imidacloprid restricted the cell migration of cardiac progenitors to primary heart field in gastrula chick embryos. A similar observation was also obtained from the cell migration assay of scratch wounds in vitro. Additionally, imidacloprid exposure negatively affected the cytoskeleton structure and expression of corresponding adhesion molecules. Taken together, these results reveal that the improper EMT, cardiac progenitor migration and differentiation are responsible for imidacloprid exposure-induced malformation of heart tube during chick embryo development.

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46	Keywords:	Imidacloprid;	chick	embryo;	heart	tube;	EMT;	cardiac	progenitor
47	migration; d	ifferentiation.							
48									

# Introduction

Organogenesis requires the precise layout of multiple cell types into a specific
three-dimensional architecture that is essential for normal organ formation. During
embryonic organ development, an obligatory process is tissue fusion, such as that of
the optic cup, palate, heart, neural tube, eyelids and body wall <sup>1, 2</sup> . Tissue fusion
appears to occur in numerous organs. Our previous study demonstrated that the
deficiency of specific transcription factors and signaling molecules could exhibit the
fusion defects in many organs, for instance, in neural tube defects <sup>3</sup> and cardiac bifida
<sup>4</sup> . As a model of organogenesis, cardiogenesis involves a series of morphogenetic
steps. In vertebrates, the heart develops from three distinct pools of cardiac
progenitors: the cardiac precursor in splanchnic mesoderm (primary and secondary
heart field), cardiac neural crest and the pro-epicardium. From the perspective of
morphological alteration, it is chronologically composed of primary heart tube fusion,
cardiac looping and accretion, cardiac septation and coronary vasculogenesis <sup>5</sup> . The
primary heart field gives rise to the major structures of the heart, including the atrias
and ventricles, while the secondary heart field contributes to the cardiac outflow tracts
<sup>6</sup> . Myocardial progenitors undergo Epithelial-Mesenchymal Transition (EMT),
proliferate, differentiation and migration into the primary heart field in the process of
heart tube formation. EMT is a morphogenetic transition process in which cells lose
their epithelial characteristics and gain mesenchymal properties underlying the
alterations of adheren junction (AJs), tight junction (TJs) and gap junction (GJs) <sup>7,8</sup> .
In the formation of primary heart fields, the precardiac cells initially migrate out of

71	the anterior primitive streak at the gastrula stage and then move symmetrically into
72	crescent location <sup>9-11</sup> . Cell migration, proliferation and differentiation are guided by its
73	micro-environment <sup>12</sup> .
74	The morphogenesis of chick cardiac looping involves four phases: pre-looping
75	phase (HH8-9); C-shaped bend (HH9+-13); S-shaped heart loop (HH14-16) and
76	primitive outflow tract formation (about 4.5 days). Within days 6-14, expansion and
77	growth of the ventricular wall benefit principally from cardiomyocyte proliferation in
78	the compact myocardium. At day 14.0, cardiac neural crest cells (CNCs) give rise to
79	the adventitia of the large veins and the coronary arteries. In this context, any
80	disruption to cardiac precursor cell migration and differentiation during cardiogenesis
81	may result in congenital heart malformations.
82	Heart development is a complex process that is tightly regulated through
83	spatio-temporal gene expression and cell-cell interaction. In previous studies of heart
84	tube assembly in the chick embryo, we have reported that fibroblast growth factor
85	(FGF) signaling, through an endoderm-derived signal, is required for regulating
86	pro-cardiac mesoderm cell migration <sup>10, 13</sup> . Additionally, bone morphogenetic protein
87	2 (BMP2) is released from the anterior endoderm and Wnt antagonists are essential
88	for precardiac mesoderm cells to differentiate into mature cardiomyocytes during
89	cardiomyogenesis <sup>14-16</sup> . Furthermore, transcription factors Nkx-2.5, GATA4,
90	myocardin and TBX5 have crucial roles in dictating morphogenesis and
91	differentiation of the heart <sup>16, 17</sup> . Vascular endothelial growth factor (VEGF) also plays
92	a vital role in the angiogenic expansion of the early network <sup>18</sup> .

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93	The neonicotinoid pesticide, imidacloprid,
94	1-((6-Chloro-3-pyridinyl)methyl)-N-nitroimidazolidinimine, has been extensively
95	used to control sucking insects, termites, soil insects on crops <sup>19</sup> and fleas on domestic
96	animals <sup>20, 21</sup> . Various products containing this chemical, including liquids, granules,
97	dusts and packages, have been sold in the US since 1994. In the EU, use of
98	imidacloprid was restricted for 2 years in 2013 because research showed a link
99	between imidacloprid and bee death (EASAC 2015, Ecosystem services, agriculture
100	and neonicotinoids). As a systemic insecticide, imidacloprid products are usually
101	sprayed on soil and leaves, and then spread to the plant's stems, leaves, fruit and
102	flowers <sup>22, 23</sup> . Imidacloprid can then penetrate into the nervous system of sucking
103	insects and combine selectively with nicotinic acetylcholine receptors (nAChR),
104	producing toxic effects <sup>24</sup> . When insects consume plants treated with imidacloprid
105	products, their nervous systems are damaged leading to death. Due to steric conditions
106	at the nAChR, imidacloprid has much lower toxicity to mammals. However, humans
107	can be exposed to imidacloprid products via skin/eye contact or through consumption
108	or inhalation when handling the pesticide or an animal recently exposed to
109	imidacloprid. The toxicity of imidacloprid in human adults is due to disruption of
110	nervous system signal transduction <sup>25</sup> . Once humans are exposed, imidacloprid
111	products can cross the lining of the intestine and be transported to the whole body
112	through circulation of the blood. However, little is known about its potential toxic
113	effects on early embryo development apart from a few reports on human health such
114	as reproductive ability. Currently, increasing attention is being paid to the toxic effects

of pesticides on embryo development, including cardiovascular system. Unfortunately, as yet there is no direct evidence of toxicological effects on cardiogenesis or corresponding mechanisms. In this study, a chick embryo model <sup>26</sup> has been used to investigate whether or not imidacloprid could affect cardiogenesis and, if so, to elucidate the underlying cellular and molecular mechanism.

#### Materials and methods

### Chick manipulations

Fertilized leghorn eggs were acquired from the Avian Farm of South China Agriculture University (Guangzhou, China). Two approaches were employed to carry out the imidacloprid exposure in this study. The imidacloprid powder was dissolved in dimethyl sulfoxide (DMSO), 0.1% DMSO was used as control to observe the potential effect of the solvent.

For imidacloprid exposure at the early embryonic stage, Hamburger-Hamilton (HH) stage 0 chick embryos from fertilized eggs were incubated with either 0.1% DMSO (control) or 500  $\mu$ M imidacloprid <sup>27</sup> in early chick (EC) culture medium in a humidified incubator (Yiheng Instruments, Shanghai, China) at 38°C and 70% humidity until the chick embryos developed to the HH10 stage. Alternatively, 500  $\mu$ M imidacloprid was directly applied to one side of the gastrula-stage embryos, with the other side being exposed to 0.1% DMSO as a control.

For imidacloprid exposure at a later embryonic stage, HH4 chick embryos were exposed to either 0.1% DMSO (control) or 500  $\mu$ M imidacloprid through injection

into windowed eggs *in vivo* and then further incubated for 4.5 days and 14 days. The experiments were performed in triplicate with 20 eggs assigned to each group, and surviving embryos were harvested for further assessment.

## In situ hybridization

Whole-mount *in situ* hybridization of chick embryos was performed according to a standard *in situ* hybridization protocol <sup>28</sup>. Briefly, digoxigenin-labeled probes were synthesized for VMHC <sup>29</sup>, GATA5 <sup>30</sup>, BMP2 and NKX2.5 (supplied by Dr. Thomas M. Schultheiss). The whole-mount stained embryos were photographed and then frozen sections prepared on a cryostat microtome (LeicaCM1900) at a thickness of 15–20 mm.

# Immunofluorescent staining

Chick embryos were harvested at the end of the experiment and fixed overnight in 4% paraformaldehyde at 4°C. Whole-mount embryos were immunofluorescently stained using MF20 (1:500, DSHB, USA), E-cadherin (1:50, BD Transduction Laboratories, USA), Laminin (1:100, DSHB, USA) antibodies. Briefly, the fixed embryos were incubated with these primary antibodies at 4°C overnight on a rocker. Following extensive washing, the embryos were incubated with the appropriate anti-mouse IgG conjugated to Alexa Fluor 488 or anti-rabbit IgG conjugated to Alexa Fluor 555 (1:1000, Invitrogen, USA), overnight at 4°C on a rocker. All embryos were finally counterstained with DAPI (1:1000, Invitrogen, USA) at room temperature for 1 hour.

# RNA isolation and semiquantitative RT-PCR

Total RNA was isolated from HH4, HH8 chick embryos using a Trizol kit
(Invitrogen, USA) according to the manufacturer's instructions. First-strand cDNA
was synthesized to a final volume of 25µl using SuperScript RIII first-strand
(Invitrogen, USA). Following reverse transcription, PCR amplification of the cDNA
was performed as described previously. The primers used for RT-PCR are provided in
the Figure S3. The PCR reactions were performed on a Bio-Rad S1000TM Thermal
cycler (Bio-Rad, USA). The final reaction volume was 50 $\mu$ l composed of 1 $\mu$ l of
first-strand cDNA, 25 $\mu M$ forward primer, 25 $\mu M$ reverse primer, 10 $\mu l$
PrimeSTARTM Buffer (Mg <sup>2+</sup> plus), 4µl dNTPs Mixture (TaKaRa, Japan), 0.5 µl
PrimeSTARTM HS DNA Polymerase (2.5U/µl TaKaRa, Japan) and RNase-free water.
The cDNA was amplified for 30 cycles. One round of amplification was performed at
94°C for 30 s, 30 s at 58°C, and 30 s at 72°C. The PCR products (20 μl) were resolved
using 1% agarose gels (Biowest, Spain) in 1× TAE buffer (0.04 M Trisacetate and
0.001 MEDTA) and 10,000x GeneGreen Nucleic Acid Dye (Tiangen, China) solution.
The resolved products were visualized using a transilluminator (Syngene, UK) and
photographs captured using a computer-assisted gel documentation system (Syngene).
The housekeeping gene GAPDH was run in parallel to confirm that equal amounts of
RNA were used in each reaction. The ratio between intensity of the fluorescently
stained bands corresponding to genes and GAPDH was calculated to quantify the
level of the transcripts for those genes mRNAs. The RT-PCR result was representative
of three independent experiments.

# 180 Cell trace with Dil

Carbocyanine dye 1, 1V-dioctadecyl-3, 3, 3V, 3V-tetramethylindocarbocyanine
perchlorate (DiI, Molecular Probes, Inc.) was used to label small groups of primitive
streak cells. A 2.5% stock solution of DiI was diluted in ethanol, 1:10 in 0.3 M
sucrose, and injected into the anterior primitive streak of HH3 chick embryo by air
pressure through a micropipette, which was pulled from a 1 mm glass capillary in a
vertical micropipette puller (WD-2, Chengdu Instrument Company). In general, each
labeled tissue in the anterior primitive streak contained approximately 10–30 cells.

#### Cell lines and culture

The H9c2 rat cardiac myoblast cell line was obtained from ATCC (American Type Culture Collection, CLR-1446, USA). The cells were cultured in a humidified incubator with 5% CO<sub>2</sub> at 37°C in six-well plates (1×10<sup>6</sup> cells/ml) containing DMEM (Gibco, Gaithersburg, MD, USA), supplemented with 10% fetal bovine serum (Gibco, Gaithersburg, MD, USA), and exposed to imidacloprid (500μM); 0.1% DMSO was used as a control. The cells were photographed using an inverted fluorescence microscope (Nikon, Tokyo, Japan) with NIS-Elements F3.2 software. After 12 hours incubation, immunofluorescent staining against phalloidin (F-actin, 1:1000, Invitrogen, Waltham, MA, USA) and anti-Myh7 (1:100, Proteintech, USA) was performed on the incubated H9c2 cells. A minimum of 5 images were assayed *per* treatment group. DAPI (1:1000, Invitrogen, USA) was used as counterstain.

## Migration assay

H9c2 cells were seeded in 6-well plates with DMEM (10% FBS) medium. At confluency, a wound was induced by scratching the monolayer with a 10-μl pipette tip.

The cells were then washed 3 times with sterile PBS. H9c2 cells were incubated in
serum-free DMEM medium with 500uM or 0.1% DMSO under 5% $\rm CO_2$ conditions.
Images were acquired at 12h and 24h post-scratching. At least 3 wells were analyzed
in each treatment group and the images were taken using an inverted microscope
(Nikon Eclipse Ti-U, Japan).

## Western blot

Chick embryos (HH4 and HH7) were collected and lysed with CytoBuster™ Protein Extraction Reagent (#71009, Novagen). The total protein concentration was established using a BCA quantification kit (BCA01, DingGuo BioTECH, CHN). Samples containing equal amounts of protein were resolved by SDS-PAGE and then transferred to PVDF membranes (Bio-Rad). The membranes were blocked with 5% Difco™ skim milk (BD) and then incubated with primary and secondary antibodies. The antibodies used were TBX5, GATA4 and GATA6 (Abcam USA), HRP-conjugated anti-mouse IgG and anti-rabbit IgG (Cell Signaling Technology, USA). All primary and secondary antibodies used were diluted to 1:1000 and 1:2000 in 5% skim milk, respectively. The protein bands of interest were visualized using an ECL kit (#34079, *Thermo* Fischer Scientific Inc.) and GeneGnome5 (Syngene). The staining intensity of the bands was determined and analyzed using Quantity One software (Bio-Rad).

# **Photography**

Following immunofluorescent staining or *in situ* hybridization, the whole-mount embryos were photographed using a stereo-fluorescent microscope (Olympus MVX10)

and associated Olympus software package Image-Pro Plus 7.0. The embryos were
sectioned into 14 $\mu$ m-thick slices using a cryostat microtome (Leica CM1900) and the
sections were then photographed with an epi-fluorescent microscope (Olympus LX51,
Leica DM 4000B) and CN4000 FISH Olympus software package.

## Data analysis

The thickness of ventricular wall and trabecular muscle and the distance of wound closure in wound healing experiments as well as the lengths of the long and short axes were all quantified with Image-Pro Plus 6.0. The cell trace with DiI experiments, Dil<sup>+</sup> cells were manually counted with Image-Pro Plus 6.0. Statistical analyses for all the experimental data was performed using a SPSS 13.0 statistical package program for Window. The data were presented as mean ± SD. Statistical significance were determined using paired T-test, independent samples T-test or one-way analysis of variance (ANOVA). \*p<0.05, \*\*p<0.01 and \*\*\*p<0.001 indicate statistically significance between control and drug-treated groups. P < 0.05 was considered to be significant.

#### Results

# Imidacloprid exposure increases cardiac malformation during chick cardiogenesis

The heart is the first functional organ in the developing embryo. There are three crucial phases in the development of heart formation: 2-, 4.5- and 14-day (Fig. 1A). To investigate the effects of excess imidacloprid exposure on heart tube formation in chick embryos, we cultured the embryos as shown in Figure S1. In the first place, we

247	found that 35% (n = 28/80), 42.5% (n = 34/80) and 50% (n = 40/80) of $500\mu M$
248	imidacloprid-treated chick embryos had died after 2, 4.5 and 14 days incubation,
249	respectively. Corresponding mortalities were only $5\%$ (n = $3/60$ ), $6.67\%$ (n = $4/60$ )
250	and $8.33\%$ (n = $5/60$ ) in the $0.1\%$ DMSO-treated chick embryos (Fig. 1B). Our results
251	showed that the growth of imidacloprid-treated embryos is slightly faster than 0.1%
252	DMSO-treated ones at 21h and, conversely, slightly delayed at 48h. (21h: DMSO =
253	$1345 \pm 74.43 \mu m$ , imidacloprid = $1629 \pm 82.45 \mu m$ , P < $0.05$ ; 48h: DMSO = $4183 \pm 10.00 \pm 10.00 \pm 10.00$
254	$45.57$ μm, imidacloprid = $3866 \pm 56.58$ μm, P < $0.001$ ; n = $40$ for each groups; Figs.
255	1C-C5, D).
256	The average number of somites in imidacloprid-treated group at 48h was about
257	10 pairs compared to 12 pairs in 0.1% DMSO-treated group (DMSO = $12.43 \pm 0.17$ ,
258	imidacloprid = $10.03 \pm 0.15$ , n = 40 for each groups, P < $0.05$ ; Fig. 1E). Next, E4.5
259	imidacloprid-treated whole embryos weights were obviously lower than 0.1%
260	DMSO-treated ones (DMSO = $0.26 \pm 0.03$ g, n = 10, imidacloprid = $0.19 \pm 0.01$ g, n =
261	34, P < 0.01; Fig. 1F). H&E staining revealed that the thicknesses of the ventricular
262	walls (DMSO = $47.52 \pm 0.95 \mu m$ , n = 10, imidacloprid = $28.85 \pm 0.72 \mu m$ , n = 14, P <
263	0.001) and the trabecular muscles were both reduced in imidacloprid-treated group
264	compared with 0.1% DMSO-treated controls (DMSO = $25.27 \pm 0.56 \mu m$ , n = $10$ ,
265	imidacloprid = $12.57 \pm 0.31 \mu m$ , n = 14, P < 0.001; Figs. 1G, G1-G2, H, H1-H2; I, J).
266	Additionally, the size and weight of imidacloprid-treated hearts were smaller and
267	lighter than those of 14-day 0.1% DMSO-treated embryos (DMSO = $0.08 \pm 0.01$ g, n
268	= 10; imidacloprid = $0.07 \pm 0.01$ g, n = 16, P < $0.05$ ; Figs. 1K, L, M). The weight of

269 whole embryo showed a similar tendency (DMSO =  $7.60 \pm 0.31$ g, n = 10, imidacloprid =  $6.03 \pm 0.29$ g. n = 16, P < 0.01; Fig. 1N). Transverse sections (Figs. 270 271 1K1, L1) and histograms established that the right ventricular wall (RV) was 272 dramatically thicker (DMSO =  $409.10 \pm 24.73 \mu m$ , n = 10, imidacloprid =  $598.20 \pm 100 \pm 100 \pm 100 \pm 100$ 273  $36.10\mu m$ . n = 16, P < 0.001; Fig. 10) whilst there was no significant difference in the 274 left ventricular wall (LV) and interventricular septum (ISV). 275 Some atypical C-looping heart tube was evident when imidacloprid-treated 276 embryos reached HH10. According to the phenotype features, we divided them into 277 four classifications: normal (Figs. 2A, B), mild (Figs. 2A1, B1), intermediate (Figs. 278 2A2, B2) and severe (Figs. 2A3, B3), and all were stained with MF20 antibody and 279 ventricular myosin heavy chain (VMHC) probe, respectively. In the 0.1% 280 DMSO-treated embryonic heart, the heart tubes are fully C-looped (normal = 100%, n 281 = 80/80), while abnormal morphological looping of heart tube occurred in the 282 imidacloprid-treated groups (normal = 13.6%, n = 8/59, mild = 39%, n = 23/59, 283 intermediate = 28.8%, n = 17/59, severe = 18.6%, n = 11/59; Fig. 2C). At stage HH10, 284 the C-shape loop of the heart tube has formed in control embryos (Fig. 2D) as 285 indicated by MF20 immunofluorescent-staining (Fig. 2E). The single cavity of the 286 heart tube was also evident in corresponding transverse sections for these 0.1% 287 DMSO-treated embryos (Figs. 2F, F1-F3). In contrast, some heart tubes of the HH10 288 imidacloprid-treated chick embryos presented in cardiac bifida (Fig. 2G), as shown in the MF20 immunofluorescently-stained heart tubes (Fig. 2H) and corresponding 289 290 transverse section of the heart tubes. The two cavities were found in the transverse

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sections of the heart tubes (Figs. 2I, I1-I3).

# Imidacloprid treatment represses cardiomyocyte differentiation

Figure 3A reveals the principal signaling pathways (Wnt, BMP&FGF and VEGF) involved in the regulation of cardiomyocyte differentiation at cardiac crescent stages (HH7-8). To explore whether imidacloprid exposure affects these crucial gene expressions of cardiomyocyte formation, we firstly exposed imidacloprid to one side of the embryos, using the other side as control. This approach has been previously described in detail <sup>31</sup>, and its advantage is in avoiding experimental artifacts due to the different velocities of embryo development. In situ hybridization results (Fig. 3B) showed that both GATA5 and Nkx2.5 expression were down-regulated on the imidacloprid-treated side, while VMHC and BMP2 expression was maintained. The results of RT-PCR showed that imidacloprid exposure increased Wnt3a expression; dramatically inhibited GATA4, TBX5, VEGFR2 and BMP4 expression, but did not affect BMP2, Fgf8 and VMHC expression (Fig. 3C). The comparisons of gene expressions are shown in Figure S.2A. The results of westren-blot showed that the imidacloprid exposure inhibited GATA4, GATA6 and TBX5 expression at protein level (Fig. 3D-E).

# Imidacloprid exposure interfered with EMT at gastrula chick embryos

Cardiac progenitor cells derived from lateral plate mesoderm cells, which were undergo EMT (Fig. 4A). During EMT, E-Cadherin down-regulation and N-cadherin up-regulation are considered to be indispensable <sup>32</sup>. Here, E-Cadherin in DMSO-treated embryos was mainly expressed in the apical side of epiblast (Figs. 4B,

B1-B1'). In contrast, expression of E-Cadherin in imidacloprid-induced embryos

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314	extended to epiblast, mesoderm and hypoblast (Figs. 4C, C1-C1'). RT-PCR showed
315	that imidacloprid treatment reduced expression of N-cadherin and increased that of
316	E-Cadherin.
317	During chick gastrulation, the earliest sign of EMT is the breakdown of BM at the
318	midline <sup>33-35</sup> . Compared to 0.1% DMSO-treated embryos (Figs. 4D, D1-D1'),
319	imidacloprid treatment shortened the midline distance (DMSO = 241.80 $\pm$ 13.99 $\mu m,n$
320	= 10, imidacloprid = 170.50 $\pm$ 7.60 $\mu$ m, n = 10, P < 0.01; Figs. 4E, E1-E1', F),
321	implying that EMT was delayed. RT-PCR data (Fig. 4G) showed no significant
322	difference between the expression of RhoA between DMSO and imidacloprid groups.
323	Imidacloprid treatment reduced the expression of P120, $\beta$ -catenin, CX43 and
324	claudin12; increased the expression of Vinculin, Par3 and occluding, but had no effect
325	on expression of AJs and TJs, including Wnt3a, Claudin-1, ZO-1 and $\alpha\text{-actin.}$ As a
326	result, it is concluded that imidacloprid treatment induced delayed EMT during
327	cardiogenesis in gastrula chick embryo. The comparisons of gene expressions are
328	shown in Figure S. 2B and C. The results of westren-blot showed that the expression
329	imidacloprid exposure down regulated N-cadherin, but up regulated E-cadherin at
330	protein level (Figs. 4H-I).
331	Imidacloprid inhibited the migration of cardiac progenitor cells
332	Cardiac progenitor cells are the resources of the heart tube and migrate bilaterally

in the lateral plate mesoderm to eventully form the cardiac crescent  $^{9, \ I0}$ . To follow the

migration trajectory of cardiac progenitor cells, DiI dye was injected into anterior

primitive streaks in HH3 chick embryos as shown in Figs. 5A, 5B. The embryos were
then exposed and cultured on either 0.1% DMSO (control) on both sides (Fig. 5A) or
with imidacloprid on one side (Fig. 5B). The photographs were taken after 9-hour and
20-hour incubations. The results showed that the Dil <sup>+</sup> mesoderm cells in the control
group migrated symmetrically at bilateral sides of embryos (n = $18$ , P > $0.05$ ; Figs.
5C-E, C1-E1, F), while many fewer Dil <sup>+</sup> mesoderm cells were observed after 9- and
20-hour incubations at the side of imidacloprid-treatment compared to the control
(DMSO = $91.00 \pm 1.38$ , imidacloprid = $43.38 \pm 1.45$ , n = $18$ , P < $0.001$ ; Figs. 5G-I,
G1-I1, J-K) This difference in Dil <sup>+</sup> cardiac progenitor cell migration clearly suggests
that imidacloprid exposure restrained the cell migration of cardiac precursors towards
the site of heart tube formation.
Imidacloprid exposure suppressed the migration, polarization, and protrusion
Imidacloprid exposure suppressed the migration, polarization, and protrusion formation of cardiac cells in vitro.
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formation of cardiac cells in vitro.  To examine the behavior of treated cells, we used H9c2 cells cultured in vitro in presence of imidacloprid. The scratch-wound assay showed that imidacloprid exposure inhibited H9c2 cells migration, as reflected in the extent of "wound" closure
formation of cardiac cells in vitro.  To examine the behavior of treated cells, we used H9c2 cells cultured in vitro in presence of imidacloprid. The scratch-wound assay showed that imidacloprid exposure inhibited H9c2 cells migration, as reflected in the extent of "wound" closure after 24h incubation from the 0.1% DMSO and imidacloprid-treated groups
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357	microscopy demonstrated that compared to 0.1% DMSO exposure (Figs. 6E-F, I)
358	imidacloprid exposure (Figs. 6G-H, J) caused a loss of cell polarization. To quantify
359	this effect, the ratios of long to short axes of cells exposed to either DMSO or
360	imidacloprid were calculated.
361	Elongation of cells exposed to imidacloprid was significantly less than that of
362	0.1% DMSO-treated control cells (DMSO = 3.13 $\pm$ 0.24 $\mu$ m, n = 25, imidacloprid =
363	$2.31 \pm 0.11 \mu m$ , n = 25, P < 0.01; Fig. 6K). More cell protrusions occurred in the
364	majority of cells exposed to 0.1% DMSO compared to those treated with imidacloprid
365	(DMSO = $85.69 \pm 3.19$ , imidacloprid = $59.79 \pm 2.89$ , n = 10 for each group, P < $0.01$ ;
366	Fig. 6L). In addition, the fluorescence intensities of Myh7 were determined (DMSO =
367	$188.50 \pm 0.94$ , n = 25, imidacloprid = $136.60 \pm 3.10$ , n = 32, P < $0.001$ ; Fig.6M).
368	RT-PCR data (Fig.6N) revealed that imidacloprid treatment reduced the expressions
369	of Vinculin, Par3, ZO-1, CX-43, Claudin-1 and α-actin, but increased the expression
370	of P120. The other tight junction gene (Claudin-12) was not affected. The
371	comparisons of gene expressions are shown in Figure S.2D. Furthermore, we also
372	detected the behavior of imidacloprid-treated chicken cardiac muscle cells <sup>37</sup> . The
373	results confirmed that imidacloprid exposure could suppress the migration,
374	polarization, and protrusion formation of cardiac cells in vitro (Figure S.4).

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# Discussion

The toxicity of imidacloprid varies greatly across species. As a neurotoxic insecticide, it has been used globally to control sucking insects in agriculture and

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animal husbandry <sup>19</sup>. Similarly, monocrotophos, an organophosphate insecticide, also has been found to greatly affect the development of zebrafish in a concentration-dependent manner 38 It has been reported that concentrations of imidacloprid in the environment was 320 µg/L near Noordwijkerhout, Nethelands, exceeding European toxicity directives, while one fifth of water samples taken in California were above the United States Environmental Protection Agency's level for invertebrates (35 µg/L for acute toxicity and 1.05 µg/L for chronic toxicity) <sup>39</sup>. Accumulation of this pesticide on plants and animals will inevitably be transferred to humans through close contacts and food contamination. A study on the biological safety of imidacloprid products is therefore particularly important <sup>27, 40</sup>. In a previous study, we conducted a concentration gradient to select the proper concentration. In our previous study, we conducted a concentration gradient to select the proper concentration. We found that mortality and ratio of malformations were both increased with the increase of the concentration <sup>41</sup>. The concentration of imidacloprid (500 µM) in this study was similar to that reported for earlier literature reports <sup>27</sup>. We considered that, for an acute toxicity experiment, the acceptable range should be less than 1000 times the environmental concentrations, and the concentration we selected here, 500  $\mu$ M (127.8 mg/L), was within this range.  $\alpha$ 7nAChR has been reported to be increased during cardiac hypertrophy in the rat <sup>42</sup>. Our previous study also found that AChR and AChE were presented in early chick embryos. We detected these expressions with acetylcholinesterase and acetylcholine receptors by RT-PCR. This work shows that expressions of both acetylcholinesterase and acetylcholine receptors

401	were inhibited by treatment with imidacloprid <sup>43</sup> . Pregnant women is a kind of
402	vulnerable groups, human embryonic development is likely to be affected by
403	cumulative toxic effects if pregnant women are exposed over the long-term to
404	imidacloprid. During embryogenesis, the heart is the first organ to be developed.
405	Severe developmental defects in the heart could cause embryonic death. Hence, it is
406	vital to determine whether or not exposure to this widely-used chemical could affect
407	development.
408	The chick embryo was selected to systemically investigate the potential toxic effect
409	of imidacloprid exposure on early heart tube formation in this study. Chick embryos
410	develop to HH10 for about 2days. Ventricular segment firstly bulge ventrally and then
411	flips to the right side. In this way, the heart fuses and a primitive C-shaped heart tube
412	is formed <sup>16</sup> . At 4.5 days, the cardiac looping process is completed <sup>44</sup> . At 14 days, the
413	expansion and growth of the ventricular wall has ended and a mature heart is
414	produced (Fig. 1A). Our results show that imidacloprid exposure significantly
415	retarded the growth of chick embryos (Fig. 1) and increased the incidence of different
416	degrees of cardiac malformations (Fig. 2). MF20, the marker of myosin II heavy chain
417	in muscles, was exploited to outline the morphology of heart tubes, and is clearly
418	expressed in the myocardium of single and complete heart tubes in 0.1%
419	DMSO-treated control (Figs. 2F, F1-F3). In contrast, the unfused cavity marked by
420	MF20 is evident in the imidacloprid-treated group (Figs. 2I, I1-I3) implying that
421	imidacloprid exposure might result in cardia bifida. Furthermore, the development of
422	ventricular wall and trabecular muscle in 4.5 days was delayed by the imidacloprid

treatment. In comparison to the reduction of cardiac volume and weight in 14 days,
the thickness of right ventricular wall was significantly increased in compensation
following imidacloprid exposure (Fig. 1O). Imidacloprid exposure-induced embryonic
mortality in the first two days is much higher than in the other two phases (Fig. 1B).
This finding also further confirms that the first two days is the crucial period for heart
tube formation. It was this period that we addressed in this study.
Morphogenesis of the heart tube during embryo development relies on a precisely
coordinated expression of cardiac-associated genes. Crescent formation mainly
requires several signal factors, including Wnt, BMP and Fgf signaling, which
coordinately control cardiomyocyte differentiation-related genes (NKX2.5,
GATA4/5/6 and T-box). Among those signal pathways, Wnt3a/ $\beta$ -catenin signal is
deemed to be a negative regulator, the others being positive <sup>16</sup> . In this study, we found
that imidacloprid exposure up-regulated Wnt3a expression and slightly
down-regulated the expression of BMP4, with not much change being observed in the
expressions of BMP2 and Fgf8. Knock-out or mutation of GATA4 and GATA5, the
zinc-finger transcription factors for cardiogenesis, leads to cardia bifida in mice $^{45}$
whilst over-expression of GATA5 induces ectopic Nkx2.5 expression. The GATA6
promoter in both mouse and chick contains functionally important Nkx2.5 binding
sites. Similarly, the murine Nkx2.5 promoter contains GATA sites that are involved in
early heart field expression <sup>46</sup> . Likewise, the unlooped heart is associated with TBX5
mutation. Furthermore, VEGFR2 and its ligand VEGF are the cardiac- and endothelial
marker at the cardiac crescent stage <sup>47</sup> . It has been observed that imidacloprid

445	exposure could result in an obvious down-regulation of VEGFR2 (Fig. 3). From the
446	results of western blot we also found the down regulaoted of GATA4, GATA6 and
447	TBX5. All these results imply that imidacloprid-treated could significantly inhibit
448	cardiomyocyte differentiation during heart tube formation.
449	It is known that cardiac crescent cells date from myocardial precursor cells initiated
450	at the anterior primitive streak of gastrula embryo. Using the Dil+ migration assay, we
451	showed that the cell migration of myocardial precursor cells was suppressed by the
452	exposure to imidacloprid (Figs. 5G-I). In comparison to the 0.1% DMSO-exposed
453	side of embryos, the less migratory Dil+ myocardial precursor cells in the
454	imidacloprid-exposed side demonstrate that imidacloprid exposure indeed interfered
455	with precardiac cell migration toward the primary heart fields. However, the
456	possibility of an influence on cell proliferation cannot be excluded.
456 457	possibility of an influence on cell proliferation cannot be excluded.  To investigate how imidacloprid affects cell migration, we employed scratch wound
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457 458 459 460	To investigate how imidacloprid affects cell migration, we employed scratch wound assay and found that exposure inhibited H9c2 cells and chicken cardiac muscle cells migration (Figs. 6B-B2, C-C2 and Fig.S4). It has been reported that cells migration properties are related to cellular cytoskeleton modulation or to relevant adherence
457 458 459 460 461	To investigate how imidacloprid affects cell migration, we employed scratch wound assay and found that exposure inhibited H9c2 cells and chicken cardiac muscle cells migration (Figs. 6B-B2, C-C2 and Fig.S4). It has been reported that cells migration properties are related to cellular cytoskeleton modulation or to relevant adherence factors <sup>48, 49</sup> . These data show that imidacloprid exposure disturbed cell internal
457 458 459 460 461 462	To investigate how imidacloprid affects cell migration, we employed scratch wound assay and found that exposure inhibited H9c2 cells and chicken cardiac muscle cells migration (Figs. 6B-B2, C-C2 and Fig.S4). It has been reported that cells migration properties are related to cellular cytoskeleton modulation or to relevant adherence factors <sup>48, 49</sup> . These data show that imidacloprid exposure disturbed cell internal structure (Fig. 6G) and reduced the number of stress fibers (Fig. 6H). Moreover, cell
457 458 459 460 461 462 463	To investigate how imidacloprid affects cell migration, we employed scratch wound assay and found that exposure inhibited H9c2 cells and chicken cardiac muscle cells migration (Figs. 6B-B2, C-C2 and Fig.S4). It has been reported that cells migration properties are related to cellular cytoskeleton modulation or to relevant adherence factors <sup>48, 49</sup> . These data show that imidacloprid exposure disturbed cell internal structure (Fig. 6G) and reduced the number of stress fibers (Fig. 6H). Moreover, cell migration also relies on cell-cell junctions, including AJs, TJs and GJs etc (Fig. 6H).

p120-catenin. Vinculin, an actin-binding protein, connects intracellular actin filaments
by forming a mixture of, for example, $\alpha$ -catenin and $\beta$ -catenin $^{50}$ . TJs located at the
top of the lateral membranes, including the claudin family and occluding, exhibit
"barrier" and "fence" functions that involve binding to intracellular ZO-1 $^{51}$ . GJs, such
as CX43, form multiple channels that allow the passage of small molecules and
electrical signals <sup>52</sup> . All the mentioned-above cell adhesion molecules were
down-regulated by imidacloprid (Fig. 6N), which suggested that this exposure
certainly interfered with cell migration and cardiac crescent formation during heart
tube formation.
Cardiac precursor cells derive from epiblast cells after undergoing EMT. EMT not
only needs to down-regulate expression of E-cadherin (required to maintain epithelial
cell contact) but also requires up-regulating the expression of N-cadherin, the
mesenchymal cell adhesion molecules. The $Wnt/\beta$ -catenin signaling pathway plays
regulatory role in the adhesion belt. Moreover, break-down of BM, marked by laminin
and the alteration of others cell-cell adhesion factors (AJs, TJs, GJs), are also very
important in EMT. In this research, imidacloprid treatment led to E-cadherin
up-regulation and N-cadherin down-regulation at mRNA and protein levels in the
gastrula chick embryos. This treatment also enhanced laminin expression but had little
influence on AJs (p120, Vinculin, Par3, $\beta$ -catenin) and GJs (CX43). These data
indicate that imidacloprid-exposure interference with EMT is achieved through
altering the relevant adhesion molecules.

In summary, these studies reveal that imidacloprid exposure negatively influenced

489	EMT, cell migration and cell differentiation in heart tube formation. Figure 7
490	summarises schematically how imidacloprid might cause these changes. But, at
491	present, the mechanisms of cardiogensis are only incompletely understood.
492	Furthermore, imidacloprid products are likely to flow into drinking water in poultry
493	farms, which may have impact on the quantity and quality of hatching eggs.
494	Thus, further experiments are required to explore the precise molecular
495	mechanism by which imidacloprid affects cardiogenesis, thereby contribute to
496	improve poultry industry.
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504	
505	Competing Financial Interest
506	The authors have declared that no competing interests exist.
507	
508	Figure legends
509	Figure 1. Imidacloprid retarded development of the chick embryos and resulted in
510	abnormal heart formation. A: The illustration shows the crucial points (2-, 4.5- and

14-day) in chick embryos heart development. B: Graph shows the mortality rate in
$0.1\%$ DMSO and $500\mu M$ imidacloprid-treated chick embryos at days 2, 4.5 and 14,
respectively. <b>C-C2</b> : Representative appearance of 0.1% DMSO-treated chick embryos
for 0- (C), 21- (C1) and 48- (C2) hs. C3-C5: Representative appearance of
imidacloprid-treated chick embryos for 0- (C3), 21- (C4) and 48- (C5) hs. <b>D</b> : Bar
chart shows the length of embryos following treatment at 0-, 21-, 48h. E: Bar chart
shows the pair numbers of somites at 48h. F: Bar chart shows the whole embryo
weight of chick embryos in E4.5. G, G1-G2: Representative appearance of the
4.5-day developing hearts in 0.1% DMSO-treated group (G), transverse section was
taken at the level indicated by dotted lines in F and stained with H&E stains (G1). The
high magnification images were taken from the sites indicated by boxed regions in G1
(G2). The black line and boxed region in G2 marked the ventricular wall and
trabecular muscle, respectively. H, H1-H2: The example shows the appearance of
4.5-day developing hearts in the imidacloprid- treated group (H), transverse section
was taken at the level indicated by dotted lines in H and stained with H&E stains (H1).
The high magnification images were taken from the sites indicated by boxed regions
in H1 (H2). The black line and boxed region in H2 dotted the ventricular wall and
trabecular muscle, respectively. I: Bar chart compares the ventricular wall thickness
of hearts. J: Bar chart compares the trabecular muscle layers. $K$ : Representative
appearance of the 14-day mature hearts in 0.1% DMSO-treated group. L: Example
shows appearance of 14-day mature hearts in the imidacloprid- treated group. K1, L1:
Transverse section was taken at the levels indicated by dashed lines in K and L. M-N:

Bar chart shows the heart weight and the whole embryo weight. **O**: The bar chart showing the thickness of ventricular wall in 14-day mature hearts. Abbreviations: LV, left ventricle; RV, right ventricle; IVS, interventricular septum. Scale bars = 2000  $\mu$ m (C, C3); 1000  $\mu$ m (C1-C2, C4-C5); 500 $\mu$ m (G-H); 300 $\mu$ m (G1-H1); 50 $\mu$ m (G2-H2); 300 $\mu$ m (K-L); 1000 $\mu$ m (K1-L1).

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Figure 2. The classification of imidacloprid exposure-induced heart malformations in gastrula chick embryos. A-A3: Representative appearances of phenotypes classification of hearts in gastrulating chick embryos immunofluorescently-stained with MF20 antibody, including normal (A), mild (A1), intermediate (A2) and severe (A3), respectively. **B-B3**: In situ hybridization shows VMHC expression in representative appearances of phenotypes classification of hearts in gastrulating chick embryos. C: Bar chart shows the rate of heart phenotype classification (%) in 0.1% DMSO- and imidacloprid-treated group. **D-E**: Representative bright-field images of 0.1% DMSO-treated HH10 embryo (D) and heart tube immunofluorescently-stained with MF20 antibody (E). F, F1-F3: F: Representative transverse sections at the levels indicated by dotted white line in E. DAPI staining is used as a counterstain in F1. F2 is the merged image. F3 is the enlarged view of boxed region in F2. G-H: Representative bright-field images of 0.1% DMSO-treated HH10 embryo (G) and heart tube immunofluorescently-stained with MF20 antibody (H). I, I1-I3: I: Representative transverse sections at the levels indicated by dotted white line in H. DAPI staining is used as a counterstain in I1. I2 is the merged image. I3 is the

555	enlarged view of boxed region in I2. Scale bars = $150 \mu m$ (A1-A4, B1-B4, E, H); $500$
556	μm (D, G); 100μm (F, F1-F3, I, I1-I3).
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558	Figure 3. Imidacloprid exposure repressed the differentiation of cardiac progenitor
559	cells. A: Overview of the signaling pathways that have been implicated into
560	cardiomyocyte formation. <b>B1-B4</b> : The embryos were incubated with 0.1% DMSO
561	(left) and imidacloprid (right) at either side until HH7 and processed for in situ
562	hybridization for GATA5 (B1), NKX2.5 (B2), VMHC (B3), BMP2 (B4). <b>B1'-B4'</b> :
563	Representative transverse sections at the levels indicated by dotted black lines in
564	B1-B4. C: RT-PCR showing the expressions at HH7 chick embryos. D: Western-bolt
565	showing the expressions at protein level in HH7 chick embryos. E: The bar chart
566	showing the comparisons of gene expressions in D. Scale bars = $200 \mu m$ (B1-B4);
567	100 μm (B1'-B4').
568	
569	Figure 4. Imidacloprid exposure interfered with EMT during chick gastrulation. A:
570	The illustration shows the EMT during chick gastrulation. B: Representative images
571	of 0.1% DMSO-treated HH4 chick embryos immunofluorescently-stained with
572	E-Cadherin. <b>B1-B1'</b> : The transverse sections at the levels indicated by dotted white
573	line in B. The section was counterstained with DAPI (B1'). E-Cadherin is expressed
574	on the apical side of epiblast of 0.1% DMSO-treated embryo (white arrow in B1'). C:
575	Representative images of imidacloprid-treated HH4 chick embryos
576	immunofluorescently-stained with E-Cadherin. C1-C1': The transverse sections at

levels indicated by dotted white line in C. The section was counterstained with L	)API
(C1'). E-Cadherin expression level was enhanced on epiblast layer, and ecc	topic
expression in the mesoderm layer following imidacloprid treatment (white arrow	vs in
C1'). <b>D</b> : Representative image of 0.1% DMSO-treated HH4 chick emb	oryos
immunofluorescently-stained for laminin. D1-D1': The transverse sections at le	evels
indicated by dotted white line in D. The section was counterstained with DAPI (I	<b>)</b> 1').
Laminin is expressed on the BM of 0.1% DMSO-treated embryo (white dotted	line
showing the gap in D1'). E: Representative image of imidacloprid-treated HH4 c	hick
embryos immunofluorescently-stained for laminin. E1-E1': The transverse section	ns at
the levels indicated by dotted white line in E. The section was counterstained	with
DAPI (E1'). Laminin is expressed on the BM of imidacloprid-treated embryo (v	vhite
dotted line showing the gap in E1'). F: Bar chart shows the gap distance of lan	ninin
(μm) with 0.1% DMSO- and imidacloprid-treated HH4 chick embryos. G: RT-	PCR
shows the expressions N-cadherinat mRNA level in the HH4 chick embryos	s. H:
Western-bolt showing the expressions at protein level in HH4 chick embryos. I:	The
bar chart showing the comparisons of gene expressions in H. Scale bars = 30	0μm
(B-E); 100μm (B1-E1, B1'-E1').	

Figure 5. Imidacloprid exposure restricted cardiac progenitor cell migration. A: The pattern of DiI-labeled cardiac progenitor cell migration following 0.1% DMSO treatment on the both sides of embryos. B: The pattern of DiI-labeled cardiac progenitor cell migration following 0.1% DMSO treatment at the left side and

imidacloprid exposure at right side of embryos. **C-E**: Fluorescence images were taken at 0- (B), 9- (C) and 20- (D) hour. Note: both sides of embryos were exposed to 0.1% DMSO. **C1-E1**: The merged images of bright-field and B-D respectively. **F**: Bar chart shows the number of cardiac precursor cells migration based on A. **G-I**: Fluorescence images were taken at 0- (G), 9- (H) and 20- (I) of incubation. The left sides of embryos were exposed to 0.1% DMSO, while the right sides were exposed to imidacloprid. **G1-I1**: The merged images of bright-field and G-I respectively. **J**: Bar chart shows the number of cardiac precursor cells migration based on F. **K**: Bar chart shows the number of embryo incidence of symmetrical migration or asymmetric migration in 0.1% DMSO- and imidacloprid groups. Scale bars = 600µm (C-E, C1-E1, G-I, G1-I1).

Figure 6. The imidacloprid exposure suppressed H9c2 cells migration, polarization and protrusion formation. A: The sketch illustrates migration of H9c2 cells as detected by the wound-healing assay. B-C: The representative images of H9c2 cells scratch test at 0-hour incubation from 0.1% DMSO-treated (B) and imidacloprid-treated (C) groups respectively. B1-C1, B2-C2: The representative images of H9c2 cells scratch test at 12-hour (B1, C1), 24-hour (B2, C2). D: The bar chart shows the percentage of wound closure (%) at 12-hour, 24-hour. E-F: Representative image of actin filaments in 0.1% DMSO -treated H9c2 cells were visualized by staining with F-actin (red), and cell nuclei were stained with DAPI (blue). White dotted lines show the long and short axes of cells. F is the enlarged view

621	of E. (The boxed region in F shows stress fiber assay in H9c2 cells). G-H:
622	Representative image of actin filaments in imidacloprid-treated H9c2 cells were
623	visualized by staining with F-actin (red), and cell nuclei were stained with DAPI
624	(blue). White dotted lines show the long and short axes of cells. H is the enlarged
625	view of G. (The boxed region in H shows stress fiber assay in H9c2 cells). I-J:
626	Representative images of 0.1% DMSO and imidacloprid-treated H9c2 cells
627	immunofluorescently-stained with Myh7, respectively. K: Bar chart showing the ratio
628	of long axis to short axis. L: Bar chart shows cells containing stress fibers (%). M:
629	Bar chart shows fluorescence intensity of Myh7 (AU). N: RT-PCR showing the
630	expressions at mRNA level in HH7 chick embryos exposed either 0.1% DMSO or
631	imidacloprid. Scale bars = $200\mu m$ (B, B1-B2, C, C1-C2); $100\mu m$ (E- J).
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633	Figure 7. Model depicting how imidacloprid exposure induced heart tube
634	malformation during chick cardiogenesis.
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642	References

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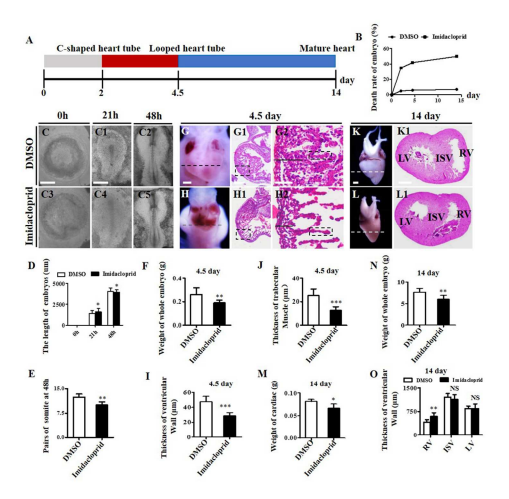


Figure 1. Imidacloprid retarded development of the chick embryos and resulted in abnormal heart formation. A: The illustration shows the crucial points (2-, 4.5- and 14-day) in chick embryos heart development. B: Graph shows the mortality rate in 0.1% DMSO and 500µM imidacloprid-treated chick embryos at days 2, 4.5 and 14, respectively. C-C2: Representative appearance of 0.1% DMSO-treated chick embryos for 0- (C), 21-(C1) and 48- (C2) hs. The embryos reached to HH10 with a C-shaped heart tube after 48 hours. C3-C5: Representative appearance of imidacloprid-treated chick embryos for 0- (C3), 21- (C4) and 48- (C5) hs. D: Bar chart shows the pair numbers of somites at 48h cultured with 0.1% DMSO and imidacloprid. E: Bar chart shows the length of 0.1% DMSO- and imidacloprid-treated embryos following treatment at 0-, 21-, 48h. F, F1-F2: Representative appearance of the 4.5-day developing hearts in 0.1% DMSO-treated group (F), transverse section was taken at the level indicated by dotted lines in F and stained with H&E stains (F1). The high magnification images were taken from the sites indicated by boxed regions in F1 (F2). The black line and boxed region in F2 marked the ventricular wall and trabecular muscle, respectively. G, G1-G2: The example shows the appearance of 4.5-day developing hearts in the imidacloprid- treated group (G), transverse section was taken at the level indicated by dotted lines in G and stained with H&E stains (G1). The high magnification images were taken from the sites indicated by boxed regions in G1 (G2). The black line and boxed region in G2 dotted the ventricular wall and trabecular muscle, respectively. H: Bar chart shows the whole embryo weight of the 0.1% DMSO- and imidacloprid-treated chick embryos in E4.5. I: Bar chart compares the ventricular wall thickness of 0.1% DMSO- and imidacloprid- treated hearts. J: Bar chart compares the trabecular muscle layers. K: Representative appearance of the 14-day mature hearts in 0.1% DMSO-treated group. L: Example shows appearance of 14-day mature hearts in the imidacloprid- treated group. K1, L1: Transverse section was taken at the levels indicated by dashed lines in K and L. M-N: Bar chart shows the whole embryo weight and the heart weight in the 14-day 0.1% DMSO- and imidaclopridtreated chick embryos. O: The bar chart showing the thickness of ventricular wall in 14-day mature hearts.

Abbreviations: LV, left ventricle; RV, right ventricle; IVS, interventricular septum. Scale bars = 2000  $\mu$ m (C1 , C4); 1000  $\mu$ m (C2-C3, C5-C6); 500 $\mu$ m (F-G); 300 $\mu$ m (F1-G1); 50 $\mu$ m (F2-G2); 300 $\mu$ m (K-L); 1000 $\mu$ m (K1-L1).

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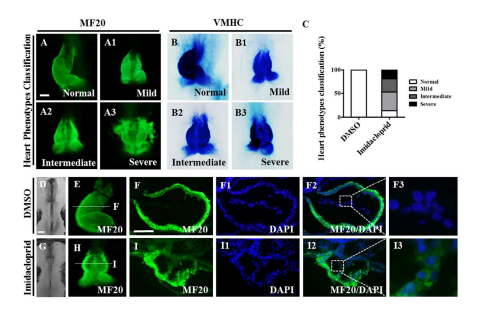


Figure 2. The classification of imidacloprid exposure-induced heart malformations in gastrula chick embryos. A-A3: Representative appearances of phenotypes classification of hearts in gastrulating chick embryos immunofluorescently-stained with MF20 antibody, including normal (A), mild (A1), intermediate (A2) and severe (A3), respectively. B-B3: In situ hybridization shows VMHC expression in representative appearances of phenotypes classification of hearts in gastrulating chick embryos, including normal (B), mild (B1), intermediate (B2) and severe (B3), respectively. C: Bar chart shows the rate of heart phenotype classification (%) in 0.1% DMSO- and imidacloprid-treated group. D-E: Representative bright-field images of 0.1% DMSO-treated HH10 embryo (D) and heart tube immunofluorescently-stained with MF20 antibody (E). F, F1-F3: F: Representative transverse sections at the levels indicated by dotted white line in E. DAPI staining is used as a counterstain in F1. F2 is the merged image of bright-field and DAPI staining. F3 is the enlarged view of boxed region in F2. G-H: Representative bright-field images of 0.1% DMSO-treated HH10 embryo (G) and heart tube immunofluorescently-stained with MF20 antibody (H). I, I1-I3: I: Representative transverse sections at the levels indicated by dotted white line in H. DAPI staining is used as a counterstain in I1. I2 is the merged image of bright-field and DAPI staining. I3 is the enlarged view of boxed region in I2. Scale bars = 150 μm (A1-A4, B1-B4, E, H); 500 μm (D, G); 100μm (F, F1-F3, I, I1-I3).

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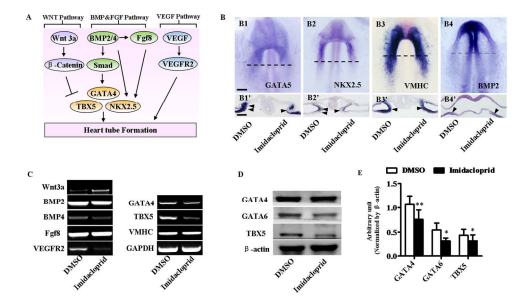


Figure 3. Imidacloprid exposure repressed the differentiation of cardiac progenitor cells. A: Overview of the signaling pathways that have been implicated into cardiomyocyte formation. B1-B4: The embryos were incubated with 0.1% DMSO (left) and imidacloprid (right) at either side until HH7 and processed for in situ hybridization for GATA5 (B1), NKX2.5 (B2), VMHC (B3), BMP2 (B4). B1'-B4': Representative transverse sections at the levels indicated by dotted black lines in B1-B4. C: RT-PCR showing the expressions at HH7 chick embryos. D: Western-bolt showing the expressions at protein level in HH7 chick embryos. E: The bar chart showing the comparisons of gene expressions in D. Scale bars = 200 μm (B1-B4); 100 μm (B1'-B4').

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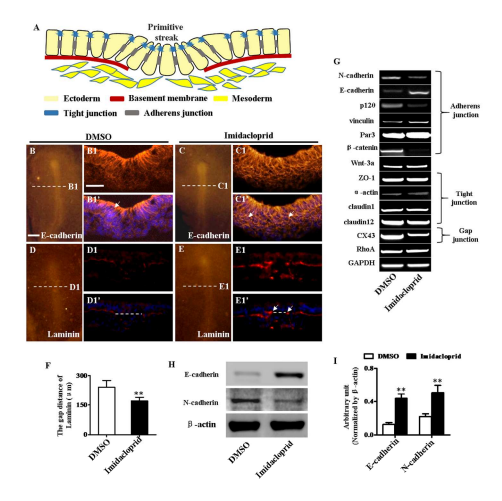


Figure 4. Imidacloprid exposure interfered with EMT during chick gastrulation. A: The illustration shows the EMT during chick gastrulation. B: Representative images of 0.1% DMSO-treated HH4 chick embryos immunofluorescently-stained with E-Cadherin. B1-B1': The transverse sections at the levels indicated by dotted white line in B. The section was counterstained with DAPI (B1'). E-Cadherin is expressed on the apical side of epiblast of 0.1% DMSO-treated embryo (white arrow in B1'). C: Representative images of imidacloprid-treated HH4 chick embryos immunofluorescently-stained with E-Cadherin. C1-C1': The transverse sections at levels indicated by dotted white line in C. The section was counterstained with DAPI (C1'). E-Cadherin expression level was enhanced on epiblast layer, and ectopic expression in the mesoderm layer following imidacloprid treatment (white arrows in C1'). D: Representative image of 0.1% DMSOtreated HH4 chick embryos immunofluorescently-stained for laminin. D1-D1': The transverse sections at levels indicated by dotted white line in D. The section was counterstained with DAPI (D1'). Laminin is expressed on the BM of 0.1% DMSO-treated embryo (white dotted line showing the gap in D1'). E: Representative image of imidacloprid-treated HH4 chick embryos immunofluorescently-stained for laminin. E1-E1': The transverse sections at the levels indicated by dotted white line in E. The section was counterstained with DAPI (E1'). Laminin is expressed on the BM of imidacloprid-treated embryo (white dotted line showing the gap in E1'). F: Bar chart shows the gap distance of laminin (µm) with 0.1% DMSOand imidacloprid-treated HH4 chick embryos. G: RT-PCR shows the expressions N-cadherinat mRNA level in the HH4 chick embryos. H: Western-bolt showing the expressions at protein level in HH4 chick embryos. I: The bar chart showing the comparisons of gene expressions in H. Scale bars = 300µm (B-E); 100µm (B1-E1, B1'-E1').

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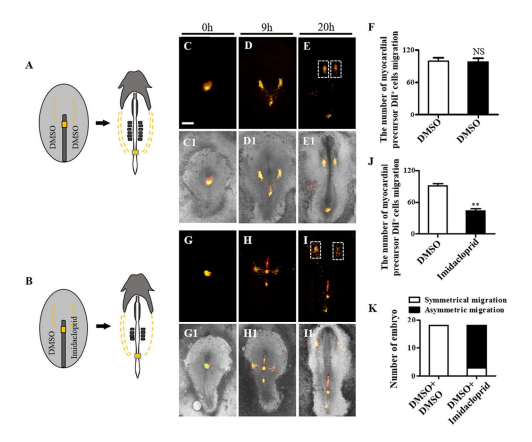


Figure 5. Imidacloprid exposure restricted cardiac progenitor cell migration. A: The pattern of DiI-labeled cardiac progenitor cell migration following 0.1% DMSO treatment on the both sides of embryos. B-D: Fluorescence images were taken at 0- (B), 9- (C) and 20- (D) hour after DiI was injected in anterior primitive streak. Note: both sides of embryos were exposed to 0.1% DMSO. B1-D1: The merged images of bright-field and B-D respectively. E: Bar chart shows the number of cardiac precursor cells migration based on A. F: The pattern of DiI-labeled cardiac progenitor cell migration following 0.1% DMSO treatment at the left side and imidacloprid exposure at right side of embryos. G-I: Fluorescence images were taken at 0- (G), 9- (H) and 20- (I) of incubation after DiI was injected in anterior primitive streak.. The left sides of embryos were exposed to 0.1% DMSO, while the right sides were exposed to imidacloprid. G1-I1: The merged images of bright-field and G-I respectively. J: Bar chart shows the number of cardiac precursor cells migration based on F. K: Bar chart shows the number of embryo incidence of symmetrical migration or asymmetric migration in 0.1% DMSO- and imidacloprid groups. Scale bars = 600μm (B-D, B1-D1, G-I, G1-I1).

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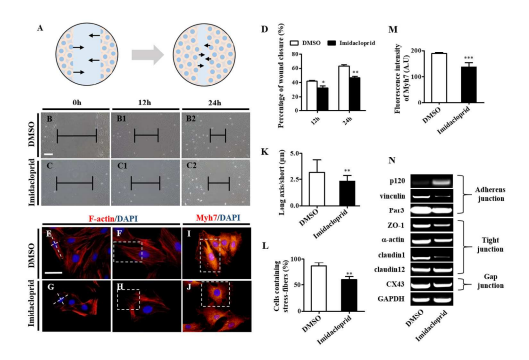
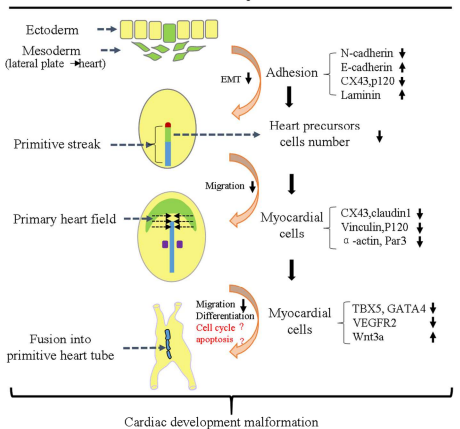


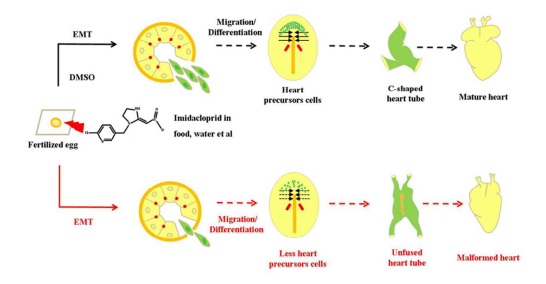
Figure 6. The imidacloprid exposure suppressed H9c2 cells migration, polarization and protrusion formation. A: The sketch illustrates migration of H9c2 cells as detected by the wound-healing assay. B-C: The representative images of H9c2 cells scratch test at 0-hour incubation from 0.1% DMSO-treated (B) and imidacloprid-treated (C) groups respectively. B1-C1, B2-C2: The representative images of H9c2 cells scratch test at 12-hour (B1, C1), 24-hour (B2, C2) incubation from 0.1% DMSO-treated (B1-B2) and imidaclopridtreated (C1-C2) groups respectively. D: The bar chart shows the percentage of wound closure (%) at 12hour, 24-hour. E-F: Representative image of actin filaments in imidacloprid-treated H9c2 cells were visualized by staining with F-actin (red), and cell nuclei were stained with DAPI (blue). White dotted lines show the long and short axes of cells. F is the enlarged view of E. (The boxed region in F shows stress fiber assay in H9c2 cells). G-H: Representative image of actin filaments in imidacloprid-treated H9c2 cells were visualized by staining with F-actin (red), and cell nuclei were stained with DAPI (blue). White dotted lines show the long and short axes of cells. H is the enlarged view of G. (The boxed region in H shows stress fiber assay in H9c2 cells). I-J: Representative images of 0.1% DMSO and imidacloprid-treated H9c2 cells immunofluorescently-stained with Myh7, respectively. K: Bar chart showing the ratio of long axis to short axis. L: Bar chart shows cells containing stress fibers (%). M: RT-PCR showing the expressions of CX43, p120, vinculin, Par3, ZO-1, a-actin, claudin1, claudin12 and RhoA at mRNA level in HH7 chick embryos exposed either 0.1% DMSO or imidacloprid. Scale bars = 200µm (B, B1-B2, C, C1-C2); 100µm (E- J).

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## Imidacloprid



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