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Source identification of airborne *Escherichia coli* of swine house surroundings using ERIC-PCR and REP-PCR

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**Abstract**

Evidence is mounting that microorganisms originating from livestock impact the air quality of the animal houses themselves and the public in the surrounding neighborhoods. The aim of this study was to develop efficient bacterial source tracking capabilities to identify sources of *Escherichia coli* aerosol pollution caused by pigs. Airborne *E. coli* were isolated from indoor air, upwind air (10 and 50 m away) and downwind air samples (10, 50, 100, 200 and 400 m away) for five swine houses using six-stage Andersen microbial samplers and Reuter-Centrifugal samplers (RCS). *E. coli* strains from pig fecal samples were also collected simultaneously. The enterobacterial repetitive intergenic consensus polymerize chain reaction (ERIC-PCR) and the repetitive extragenic palindromic (REP-PCR) approaches were used to study the genetic variability and to determine the strain relationships among *E. coli* isolated from different sites in each swine house. Results showed that 35.1% (20/57) of the bacterial DNA fingerprints from the fecal isolates matched with the corresponding strains isolated from indoor and downwind air samples (similarity ≥90%). *E. coli* strains from the indoor and downwind air samples were closely related to the *E. coli* strains isolated from feces, while those isolated from upwind air samples (swine house C) had low similarity (61–69%). Our results suggest that some strains isolated from downwind and indoor air originated in the swine feces. Effective hygienic measures should be taken in animal farms to prevent or minimize the downwind spread of microorganism aerosol.

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1. Introduction

Bioaerosols contamination caused by animals maybe a threat to both animals and public health. In particular, microorganisms or their fragments can be passively or actively released into the air in response to the changes of the hygienic situation of animal houses and meteorological conditions (Jones and Harrison, 2004). Airborne microorganisms found in swine confinement units may have a direct impact on the wellbeing of the animals in different stages of growth as well as the health of human caretakers. Many airborne pathogenic microorganisms, including viruses and bacteria, can be transmitted great distances through air flow (Brown and Hovmoller, 2002; Donaldson, 1999; Lee et al., 2007; Mims and Mims, 2004; Shinn et al., 2000). Examples include Foot-and-Mouth disease virus (Donaldson et al., 1982); severe acute respiratory syndrome virus (Eubank et al., 2004; Weiss and McMichael, 2004); influenza virus (Klontz et al., 1989; Landman and Schrier, 2004); *Bacillus anthracis* (CDC, 2003); and *Klebsiella pneumonia* (Prazmo et al., 2003). These facts illustrate the need for adequate source identification of the origins and transmission of microbial pathogens. An understanding of the origin of air pollution is paramount for assessing associated health risks as well as the actions necessary to remedy the problem.

Airborne bacterial source tracking has been studied mostly by concentration differences of the total number of aerobic bacteria (Chai et al., 1999, 2001; Kim and Kim, 2007), and the differences of antibiotic resistances of bacteria (Chai et al., 2003; Yao et al., 2007). However, tracing the origin of airborne microorganisms is difficult using culture-based approaches. Additionally, using traditional bacterial taxonomy proves problematic when differentiating between strains that share a close genetic relationship. With the development of molecular biology, genotypic techniques were developed for studying bacterial source tracking in the field. This was initially achieved in water pollution identification (Ahmed et al., 2005; Griffith et al., 2003; Indest et al., 2005; Meays et al., 2004; van Belkum et al., 1996). Molecular methods...
are able to distinguish between closely related bacteria by producing DNA fingerprints that are specific for individual strains. By the comparison of these DNA fingerprints, strains from one site can be related to strains from another site. Tracing the migration of an indicator bacterium of air pollution based on comparison of their DNA fingerprints provides invaluable information. However, this technique has not yet been applied to study the origin and transmission of bioaerosols in animal houses.

Several microorganisms have been used as indicators of fecal contamination. Among indicator bacteria, E. coli and Enterococcus spp. are most commonly used as the target organisms for bacterial source tracking in seawater (Bitton et al., 1983; Hanes and Fragala, 1987; Keswick et al., 1982; McFeters et al., 1974; Omura et al., 1982; Wait and Sobsey, 2001). E. coli are a dominant intestinal bacterial population in swine, colonizing the intestine distal to the stomach (Schierack et al., 2006). It is estimated that 1–4% of all cultivable bacteria of the colon are E. coli, and up to 105 E. coli bacteria can be detected in 1 g of feces (Sahnet et al., 2005; Selander et al., 1987). Nonpathogenic E. coli (commensal E. coli) strains are thought to maintain the physiological milieu of the gut and support digestion as well as defend against enteric pathogens.

Other E. coli strains carrying virulence genes (pathogenic E. coli), can cause severe outbreaks such as neonatal and postweaning diarrhea and edema of swine (Gyles and Fairbrother, 2004; Imberechts et al., 1992). These strains are responsible for major economic losses in pig rearing (Wieder et al., 2001). In addition, E. coli is a common microorganism in the air environment, particularly in animal houses and their surroundings (Duan et al., 2006; Northcutt et al., 2004; Zucker et al., 2000). Hojovec et al. (1977) evaluated the air quality in poultry houses using E. coli as an indicator. In this experiment, E. coli was used as indicator bacterium to study the origin of bioaerosol and its transmission to the ambient air of swine houses.

In this study, the genetic similarity of E. coli strains isolated from air samples of different sites and fecal samples in five different swine houses was characterized using the ERIC-PCR and REP-PCR. From this, a deeper understanding of the mechanisms of the occurrence and spread of airborne E. coli can be obtained.

### 2. Materials and methods

#### 2.1. Animal houses studied

Five swine houses were studied in Shandong, China, between January and October 2006. These houses were located outside of neighboring villages where there were no buildings or tall vegetation around the swine houses. Swine house A had a distance of more than 2000 m to the nearest village, where there were no crops planted at that time. Swine house B had a distance of more than 3000 m to the nearest village, where they were planting wheat at that time. Swine house C, D, and E were located to the east of the villages with distances ranging from 500 to 1000 m from the nearest village and had the same geographical characteristics. The meteorological conditions were recorded at the time of air sampling and a description of these conditions is given in Table 1.

#### 2.2. Airborne Escherichia coli

Six-stage Andersen samplers (Andersen, 1958) were used to sample the indoor air for the collection of E. coli in the swine houses. Samplers were placed near the middle of the stable about 1 m above the ground. The Reuter-Centrifugal samplers (RCS) were used to collect airborne E. coli from the outdoor air at the different sites upward (10 and 50 m away from the swine houses) and downward (50, 100, 200 and 400 m away from the swine houses). The airflow rates for the Andersen sampler and the RCS were 28.3 and 40.1 L/min, respectively. The Andersen sampler was equipped with MacConkey agar plates and operated for 1–8 min and the RCS was equipped with MacConkey agar strips and also operated for 1–8 min. At each sampling site, five air samples were collected continuously. The exposed agar plates and strips were incubated at 37 °C for 48 h. All colonies that appeared on the plates and strips were screened based on their Gram reactions using the KOH assay. Gram-negative colonies were subcultured on MacConkey agar and species identifications were made using conventional biochemical tests; the ability to ferment lactose and glucose (Kluger-Iron test); to produce hydrogen sulfide, and when in doubt, use of the API 20 E System (BioMérieux, Marcy-l’Étoile, France).

#### 2.3. Escherichia coli in feces

Ten to 15 fresh fecal samples of healthy pigs were aseptically collected from each feedlot randomly from piles on the ground and air samples were collected simultaneously. Then, the fecal samples were immediately transported to the laboratory in ice-cooled containers and analyzed within 24 h of collection. Approximately 1 g of feces was transferred to a sterilized glass homogenizer containing 9 mL of 0.9% sodium chloride. Samples were diluted and spread onto the surface of eosin methylene blue (EMB) agar plates (Tianhe, Hangzhou, China) which were incubated at 37 °C for 18–20 h. Colonies with a metallic sheen on EMB agar were picked and streaked onto MacConkey agar No. 3 (Oxoid). After overnight incubation at 37 °C, one or two typical pink colonies were selected from each MacConkey agar plate and the isolate identifications were performed as described in Section 2.2.

#### 2.4. DNA extraction

All E. coli isolates were grown in 5 mL of Luria-Bertani (LB) broth (Oxoid) with moderate shaking for 18 h at 37 °C. Then, 1.5 mL LB broth was removed and centrifuged at 10,000 g for 2 min. The cell pellet was resuspended in 100 μL of sterile double distilled water and held in boiling water for 10 min. Bacteria were then rapidly cooled by placing on ice for 5 min. This mixture was centrifuged at 12,000 g for 2 min and the supernatant removed and stored at −20 °C. These samples were used as the DNA templates for PCR analysis (Cheng et al., 2006).

#### 2.5. ERIC-PCR and REP-PCR

The PCR reaction mixture (25 μL) contained 1.5 U Taq DNA polymerase (TaKaRa), 300 ng μL−1 of each primer, 0.875 mM of each dNTP, 1× reaction buffer and 1.5 mM MgCl2 (TaKaRa). Amplification was performed in a thermocycler (Eppendorf, Germany) as follows: an initial denaturation (94 °C, 5 min) followed by 35 cycles of denaturation (94 °C, 1 min), annealing (1 min at 42 °C), and extension (65 °C, 8 min) with a single final extension (65 °C, 16 min). The reaction products were stored at 4 °C until they were electrophoresed on a 1.2–1.5% (w/v) agarose gel containing 1× TAE and 0.5 μg mL−1 ethidium bromide along with the DL2000 DNA markers (TaKaRa). The primers ERIC1, ERIC2, REP1R and REP2I were synthesized by TaKaRa (Dalian, China) and the sequences (5’–3’) are as follows (Versalovic et al., 1991, 1994):

**ERIC 1**: ATGTAAGCTCCTGGGGATTCAC; ERIC 2: AAGTAAGTGACTGGGGTGAGCG;

**REP1R**: IIIICGICGICATCIGGC; REP2I: ICGICTTATCIGGCCTAC.

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### Table 1

| Swine house | N   | Layout       | Ventilation system | Age (D)   | Inside | Outside |
|-------------|-----|--------------|--------------------|-----------|--------|---------|
|             |     |              |                    |           | T (°C) | RH (%)  | WS (m/s) | T (°C) | RH (%)  | WS (m/s) |
| A           | 450 | Close        | Mechanical         | 90–106    | 17     | 81      | 0        | 14     | 78      | 0–0.6    |
| B           | 500 | Half-close   | Natural            | 45–190    | 22     | 65      | 0        | 18     | 60      | 1.4–5.0  |
| C           | 600 | Half-close   | Natural            | 70–95     | 28     | 67      | 0        | 35     | 58      | 0–1.5    |
| D           | 360 | Half-close   | Natural            | 48–85     | 20     | 67      | 0        | 22     | 70      | 0–1.5    |
| E           | 480 | Half-close   | Natural            | 45–80     | 22     | 65      | 0        | 24     | 70      | 0–1.5    |

*Note: N = Number of animal; D = day; T = Temperature; RH = Relative Humidity; WS = Wind Speed.*
2.6. ERIC-PCR and REP-PCR fingerprints analyses

ERIC and REP-PCR fingerprints of amplified DNA fragments obtained from the agarose gel electrophoreses were recorded. The observed bands in the gels were evaluated based on the presence (coded 1) or absence (coded 0) of polymorphic fragments for the ERIC primers. Cluster analysis was performed with NTSYS-pc (Version 2.10), a numerical taxonomy and multivariate analysis software package (Rohlf, 2000), based on Dice’s similarity coefficient \((S)\) with a 1% position tolerance and the unweighted pair group method using arithmetic averages (UPGMA). In addition, each isolate was considered as an operational taxonomic unit (OTU). In order to reduce the number of OTUs in the dendrogram, to facilitate interpretation, isolates of >90% similarity were treated as a single isolate (Borges et al., 2003).

3. Results

The number of isolates recovered and the number examined from each site (including fecal samples) at each swine house was presented in Table 2. Table 2 showed that a total of 60 fecal samples were examined, and 57 \(E. coli\) strains were detected in this experiment. At the same time, 41 \(E. coli\) strains from indoor air and 19 \(E. coli\) strains from downwind air were detected. Only 3 \(E. coli\) strains from upwind air were detected at swine house C (Table 2).

The electrophoretic profiles of the ERIC-PCR products were determined for the \(E. coli\) strains isolated from indoor air, outdoor air and fecal samples. The data matrices based on the DNA fragments and the dendrogram using the NTSYS-pc software were constructed, grouping all \(E. coli\) isolates obtained in this study into different clusters or branches based on the ERIC-PCR pattern similarity. For best comparision and resolution, the ERIC DNA fragments from the isolates obtained from the same animal house were amplified in one PCR reaction and loaded on the same gel.

According to the UPGMA dendrogram (Figs. 1-5), each swine house contained fecal-obtained \(E. coli\) strains that shared 100% similarity with the corresponding air-obtained \(E. coli\) strains. Additional air sampled strains obtained either upwind, downwind, or indoor had the similarity ranged from 100% to 58% (swine house A); to 43% (swine house B); to 61% (swine house C); to 52% (swine house D); to 50% (swine house E) with the corresponding strains isolated from fecal samples. The number of \(E. coli\) strains isolated from indoor air which shared 100% similarity with fecal strains was different in 5 swine houses (Table 3).

3.1. The similarity among \(E. coli\) strains isolated from indoor air and fecal samples

As shown in the UPGMA dendrograms, 33.3% (14/42) of \(E. coli\) strains (Table 3) isolated from indoor air samples shared 100% similarity with corresponding strains isolated from fecal samples in the five swine houses. In swine house A, strains indoor-3 and -4 shared 100% similarity with feces-6, while the same was observed for indoor-1 and feces-9 (swine house B); indoor-1,-11 and feces-1 (swine house C); indoor-8 and feces-3,-10 (swine house D); indoor-9 and feces-5 (swine house D); indoor-3 and feces-5 (swine house E); (Figs. 1-5). In addition, 23.8% (10/42) of \(E. coli\) strains isolated from indoor air samples had >90% but <100% similarity with the fecal strains. In swine house A, indoor-1 and feces-6 shared 93% similarity; in swine house B, indoor-3,-6 and feces-9 shared 92% similarity as did indoor-5 and feces-10; in swine house C, over 90% similarity existed among indoor-2,-4 and feces-1,-3,-6,-10; in swine house D indoor-2 and feces-12 shared 93% similarity; in swine house E, indoor-4 and feces-5 had 94% similarity (Figs. 1-5).

Of the strains isolated from indoor air samples, 50.0% (21/42) had <90% similarity with strains isolated from fecal samples in

![Fig. 1. ERIC-PCR and REP-PCR fingerprints of 15 E. coli strains in swine house A. The dendrogram was constructed using UPGMA with a 1% band tolerance (feces-1 means the first E. coli strain isolated from feces; indoor-2 means the second E. coli strain isolated from indoor air; upwind50m-1 means the first E. coli strain isolated from upwind at 50 m; downwind100m-1 means the first E. coli strain isolated from downwind at 100 m).](image1)

![Fig. 2. ERIC-PCR and REP-PCR fingerprints of 23 E. coli strains in swine house B. The dendrogram was constructed using UPGMA with a 1% band tolerance (feces-1 means the first E. coli strain isolated from feces; indoor-2 means the second E. coli strain isolated from indoor air; upwind50m-1 means the first E. coli strain isolated from upwind at 50 m; downwind100m-1 means the first E. coli strain isolated from downwind at 100 m).](image2)

| Site           | Swine house | A (R/E) | B (R/E) | C (R/E) | D (R/E) | E (R/E) |
|----------------|-------------|---------|---------|---------|---------|---------|
| Fecal          |             | 10/10   | 10/10   | 14/15   | 12/15   | 11/10   |
| Upwind 50 m    |             | 0/5     | 0/5     | 1/5     | 0/5     | 0/5     |
| Upwind 10 m    |             | 0/5     | 0/5     | 2/5     | 0/5     | 0/5     |
| Indoor air     |             | 5/5     | 8/5     | 12/5    | 9/5     | 7/5     |
| Downwind 10 m  |             | 0/5     | 2/5     | 4/5     | 2/5     | 3/5     |
| Downwind 50 m  |             | 0/5     | 2/5     | 3/5     | 2/5     | 1/5     |
| Downwind 100 m |             | 0/5     | 1/5     | 1/5     | 0/5     | 0/5     |
| Downwind 200 m |             | 0/5     | 0/5     | 0/5     | 0/5     | 0/5     |
| Downwind 400 m |             | 0/5     | 0/5     | 0/5     | 0/5     | 0/5     |

Note: R/E = The number of isolates recovered/the number examined.
the five swine houses. For instance, in swine house A indoor-5 and feces-8, 9 shared 79% similarity; in swine house B indoor-8 and feces-9 had 79% similarity. All isolates with >90% similarity were treated as independent strains.

3.2. The similarity among E. coli strains isolated from downwind air samples and indoor air or fecal samples

A total of 36.8% (7/19) E. coli strains (Table 3) isolated from downwind air samples at different sampling sites had 100% similarity with corresponding strains isolated from indoor air or fecal samples in the five swine houses. For instance, in swine house B downwind10m-1 and indoor-8 had 100% similarity as did downwind10m-2 and feces-5, downwind50m-2 and feces-6, downwind10m-3 and feces-9 (swine house C); downwind10m-1, feces-8 and indoor-7 (swine house D); downwind10m-3 and feces-6 (swine house E) (Figs. 2–5). In addition, 15.8% (3/19) E. coli strains isolated from downwind air samples at different sampling sites had >90% but <100% similarity with corresponding strains isolated from indoor air or fecal samples. For example, in swine house B, downwind50m-1 and indoor-8 shared 92% similarity; in swine house C downwind100m-1 and feces-5 shared 94% similarity; in swine house E downwind10m-1, indoor-3 and feces-5 had 92% similarity (Figs. 2, 3, 5). Isolates with >90% similarity were treated as the same strain (Borges et al., 2003).

In addition, 47.4% (9/19) E. coli strains (Table 3) isolated from downwind air samples had <90% similarity with corresponding strains isolated from indoor air or fecal samples in the five swine houses. In swine house B downwind10m-1 and indoor-8 had 100% similarity as did downwind10m-2 and feces-5, downwind50m-2 and feces-6, downwind10m-3 and feces-9 (swine house C); downwind10m-1, feces-8 and indoor-7 (swine house D); downwind10m-3 and feces-6 (swine house E) (Figs. 2–5). In addition, 15.8% (3/19) E. coli strains isolated from downwind air samples at different sampling sites had >90% but <100% similarity with corresponding strains isolated from indoor air or fecal samples. For example, in swine house B, downwind50m-1 and indoor-8 shared 92% similarity; in swine house C downwind100m-1 and feces-5 shared 94% similarity; in swine house E downwind10m-1, indoor-3 and feces-5 had 92% similarity (Figs. 2, 3, 5). Isolates with >90% similarity were treated as the same strain (Borges et al., 2003).

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3.3. The similarity among E. coli strains isolated from upwind air samples and indoor air or fecal samples

Airborne E. coli strains were isolated from upwind air samples only for swine house C and all three of these strains (upwind10m-1, 2 and upwind50m-1) had low similarity (61–69%) with corresponding strains isolated from indoor air or fecal samples (Fig. 3, Table 3).

4. Discussion

An effective method for bacterial source tracking is one that can correctly identify the origin of environmental isolates; has the
samples had a far genetic distance (\(o\) 100% 90–100% similarity) with strains isolated from indoor air or fecal samples or outdoor air samples based on the ERIC-PCR and REP-PCR methods in 5 swine houses. More specifically, 48.8% of strains isolated from indoor air samples had a far genetic distance (90% similarity) with strains isolated from indoor air or fecal samples or outdoor air samples. This observation might result from the aerosolization of microorganisms originating from five swine houses. As illustrated in Figs. 1–5, results indicated that some strains isolated from downwind sites did not originate from the swine stables. It is obvious that outdoor air can be easily affected by anthropogenic activities (Lighthart, 1984) and environmental factors.

**Airborne** *E. coli* was not collected in upwind air samples from four of the swine houses (A, B, D and E) or 200 m downwind of all five swine houses. This was likely due to the following reasons: (i) the concentrations of airborne *E. coli* were very low in these sampling sites; (ii) environmental factors such as relative humidity, temperature, ultraviolet (UV) radiation, oxygen content, specific ions, various pollutants and air-associated factors influenced the ability of microorganisms to survive in the air (Herrmann et al., 2007); (iii) the aerosolization of microorganisms and sampling stress lead to a loss of culture viability (Marthi et al., 1999; Stewart et al., 1995).

Based on these findings, it can be concluded that the use of ERIC-PCR and REP-PCR are beneficial, and these approaches have shown to be useful for assessing the genetic diversity of isolates from different geographic locations (Mehta et al., 2002). Moreover, the consumables and labor costs for ERIC-PCR and REP-PCR analyses are significantly lower than those for other genotyping methods (Olive and Bean, 1999).

Taken together, by using the accurate ERIC-PCR and REP-PCR technologies, *E. coli* in feces was found aerosolized and spread to outdoor air, especially to the downwind air of the swine houses via air exchange. Airborne microbes released from swine farms may pose health risks to exposed animals and human beings, especially farm workers (Donham et al., 1986). However, further studies are needed to determine if these microbial contaminants actually cause measurable human or animal health effects.

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**Table 3**

Similarity among *E. coli* strains isolated from indoor air and fecal samples or outdoor air samples based on the ERIC-PCR and REP-PCR methods in 5 swine houses.

| Stable | The number of strains | Indoor air with fecal samples | Indoor air or fecal samples with downwind samples | Indoor air with upwind samples |
|--------|-----------------------|-------------------------------|-----------------------------------------------|-------------------------------|
|        |                       | 100% 90–100% <90%             | 100% 90–100% <90%                               | 100% 90–100% <90%             |
| A      | 2/5                   | 1/5                           | –                                             | –                             |
| B      | 1/8                   | 3/8                           | 4/8                                           | 1/5                           |
| C      | 7/12                  | 0/12                          | 5/12                                          | 4/8                           |
| D      | 2/9                   | 1/9                           | 6/9                                           | 1/2                           |
| E      | 2/8                   | 2/8                           | 4/8                                           | 1/4                           |

**Note:** ‘–’ = *E. coli* strains were not isolated in these sites.

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**ERIC also known as intergenic repeat units is another family of prokaryotic repetitive sequences. In *E. coli* there are 30–50 copies of ERIC sequences, and ERIC units tend to occur singly, including a potential stem-loop structure, and are located in non-coding regions (Hultén et al., 1991; Martin et al., 1992; Sharples and Lloyd, 1990). REP sequences are present from 500 to 1000 base pairs in length and are found in clusters in which successive copies are found in alternate orientations (Gilton et al., 1984; Martin et al., 1992; Stern et al., 1984). And REP-PCR is a widely used method for DNA typing analysis and has been shown to successfully differentiate strains of Bartonella (Rodriguez-Barradas et al., 1995), Citrobacter diversus (Woods et al., 1992), Enterobacter aerogene (Georghiou et al., 1995), Burkholderia cepacia (Hamill et al., 1995), Salmonella enterica (Johnson and Calbots, 2000) and *E. coli* (Dombek et al., 2000). So, the ERIC-PCR and REP-PCR techniques have been devised for characterization of bacteria and are widely employed to distinguish species, strains and serotypes (Chansrirornchai et al., 2001; de Moura et al., 2001; Lupski and Weinstock, 1992; van Belkum, 1994; Versalovic et al., 1994). McLellan et al. (2003) found that ERIC-PCR and REP-PCR yielded comparable results using *E. coli* as the target organism. In this experiment, our results showed that the discriminatory power yielded by the techniques of ERIC-PCR and REP-PCR was very well to differentiate between strains isolated from different sites in each swine house.

In the present study, we analyzed 120 *E. coli* isolates originating from five swine houses. As illustrated in Figs. 1–5, the grouping of REP-PCR fingerprints was highly similar with those obtained from ERIC-PCR. This showed that the two approaches yield an analogous taxonomic resolution, suggesting that the two approaches can be fully applied for the study of the transmission of bioaerosols from animal houses to their environments. From the results, we can see that 48.8% *E. coli* strains isolated from indoor air samples had >90% similarity with corresponding strains isolated from fecal samples in each swine house. That is, the fecal samples could be aerosolized and spread to the indoor air. In contrast, however, 51.2% of *E. coli* isolated from indoor air samples had a far genetic distance (<90% similarity) from the fecal samples. This observation might result from the following reasons: (i) the strains isolated from the indoor air originated from other sources (e.g. feed, water, etc.); (ii) these strains are present in the fecal samples, but were not isolated from the fecal samples and could not be compared.

Similarly, 52.6% (10/19) of strains isolated from downwind air samples had >90% similarity with strains isolated from indoor air and/or fecal samples, indicating that the strains in the indoor air could spread to the outdoor air, especially the downwind air, via air exchange. Some strains isolated from downwind (47.4%) however had a <90% similarity with strains corresponding from the indoor air or fecal samples in swine houses B–C, while no *E. coli* strains were isolated from downwind of house A. These results indicated that some strains isolated from downwind sites did not originate from the swine stables. It is obvious that outdoor air can be easily affected by anthropogenic activities (Lighthart, 1984) and environmental factors.

**ERIC and REP-PCR are widely employed to distinguish species, strains and serotypes** (Chansrirornchai et al., 2001; de Moura et al., 2001; Lupski and Weinstock, 1992; van Belkum, 1994; Versalovic et al., 1994). McLellan et al. (2003) found that ERIC-PCR and REP-PCR yielded comparable results using *E. coli* as the target organism. In this experiment, our results showed that the discriminatory power yielded by the techniques of ERIC-PCR and REP-PCR was very well to differentiate between strains isolated from different sites in each swine house.

In the present study, we analyzed 120 *E. coli* isolates originating from five swine houses. As illustrated in Figs. 1–5, the grouping of REP-PCR fingerprints was highly similar with those obtained from ERIC-PCR. This showed that the two approaches yield an analogous taxonomic resolution, suggesting that the two approaches can be fully applied for the study of the transmission of bioaerosols from animal houses to their environments. From the results, we can see that 48.8% *E. coli* strains isolated from indoor air samples had >90% similarity with corresponding strains isolated from fecal samples in each swine house. That is, the fecal samples could be aerosolized and spread to the indoor air. In contrast, however, 51.2% of *E. coli* isolated from indoor air samples had a far genetic distance (<90% similarity) from the fecal samples. This observation might result from the following reasons: (i) the strains isolated from the indoor air originated from other sources (e.g. feed, water, etc.); (ii) these strains are present in the fecal samples, but were not isolated from the fecal samples and could not be compared.
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