Tracing of the Bile-Chemotactic Migration of Juvenile *Clonorchis sinensis* in Rabbits by PET-CT

Tae Im Kim1, Won Gi Yoo1,2, Byung Kook Kwak3, Ju-Won Seok4, Sung-Jong Hong1*

1 Department of Medical Environmental Biology, Chung-Ang University College of Medicine, Seoul, Republic of Korea, 2 Division of Malaria and Parasitic Diseases, National Institute of Health, Korea Centers for Disease Control and Prevention, Osong, Chungbuk, Republic of Korea, 3 Department of Radiology, Chung-Ang University College of Medicine, Dongjak-gu, Seoul, Republic of Korea, 4 Department of Nuclear Medicine, Chung-Ang University College of Medicine, Dongjak-gu, Seoul, Republic of Korea

**Abstract**

**Background:** Adult *Clonorchis sinensis* live in the bile duct and cause clonorchiasis. It is known that the *C. sinensis* metacercariae excyst in the duodenum and migrate up to the bile duct through the common bile duct. However, no direct evidence is available on the *in vivo* migration of newly excysted *C. sinensis* juveniles (CsNEJs). Advanced imaging technologies now allow the *in vivo* migration and localization to be visualized. In the present study, we sought to determine how sensitively CsNEJs respond to bile and how fast they migrate to the intrahepatic bile duct using PET-CT.

**Methodology/Principal Findings:** CsNEJs were radiolabeled with 18F-fluorodeoxyglucose (18F-FDG). Rabbits with a gallbladder contraction response to cholecystokinin-8 (CCK-8) injection were pre-screened using cholecintigraphy. In these rabbits, gallbladders contracted by 50% in volume at an average of 11.5 min post-injection. The four rabbits examined were kept anesthetized and a catheter inserted into the mid duodenum. Gallbladder contraction was stimulated by injecting CCK-8 (20 ng/kg every minute) over the experiment. Anatomical images were acquired by CT initially and dynamic PET was kept anesthetized and a catheter inserted into the mid duodenum. Gallbladder contraction was stimulated by injecting CCK-8 (20 ng/kg every minute) over the experiment. Anatomical images were acquired by CT initially and dynamic PET was then carried out for 90 min with a 3-min acquisition per frame. Twelve minutes after CCK-8 injection, about 3,000 18F-FDG-labeled CsNEJs were inoculated into the mid duodenum through the catheter. Photon signals were detected in the liver 7–9 min after CsNEJs inoculation, and these then increased in the whole liver with stronger intensity in the central area, presenting that the CsNEJs were arriving at the intrahepatic bile ducts.

**Conclusion:** In the duodenum, CsNEJs immediately sense bile and migrate quickly with bile-chemotaxis to reach the intrahepatic bile ducts by way of the ampulla of Vater.

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* E-mail: hongsj@cau.ac.kr

**Introduction**

Human *Clonorchis sinensis* infections are endemic in East Asia countries, such as China, Vietnam, and Korea, where 15–20 million people are estimated to be infected [1]. In South Korea, clonorchiasis is currently the most prevalent parasitic infection and estimated to infect 1.3 million people [2]. *C. sinensis* infected patients suffer from abdominal pain, hepatomegaly, obstructive jaundice, indigestion, and complications of cholecystitis, cholangitis, and cholangiocarcinoma [3,4]. Furthermore, recently, *C. sinensis* was categorized as a Group 1 biological carcinogen by the International Agency for Research on Cancer [5].

Humans are the final host and become infected by eating freshwater fish containing *C. sinensis* metacercariae. Ingested metacercariae excyst in the duodenum due to trypsin stimulation [6], and the newly excysted *C. sinensis* juveniles (CsNEJs) migrate to the intrahepatic bile duct. The migration route of CsNEJs has been previously examined in experimental animals. In rabbit experiments, the common bile duct was first ligated surgically then *C. sinensis* metacercariae were administered to the rabbits through a gastric tube. One month later, adult *C. sinensis* were searched for in the bile ducts, but were not found. Based on this finding it was suggested that CsNEJs migrate through the common bile duct to the intrahepatic bile ducts [7], and this has been taken to be the migration route of *C. sinensis* in mammalian hosts [3].

Parasites such as *C. sinensis* have specific *in vivo* migration routes in their hosts, which could be targeted for development of therapeutic and preventive interventions against parasitic diseases. Furthermore, *in vivo* imaging technologies have been recently developed for the clinical diagnoses of a wide range of diseases, and these techniques have a potential to monitor the movements of CsNEJs.

Molecular imaging has emerged as a discipline at the intersection of molecular biology and *in vivo* imaging. It enables cellular functions to be visualized and molecular processes to be followed in living organisms in a non-invasive manner. Recently, studies on the visualization of live parasite in hosts have been conducted. Using transgenic *Plasmodium* parasites, pre-erythrocytic development was visualized; *Plasmodium* sporozoites entered hepatic cells, developed in a large schizont, and released merozoites in liver [8,9]. However, these techniques are not applicable to trematodes, because stable transgenic flukes are difficult to be generated.
In mammalian hosts, adult forms of trematodes consume large amounts of glucose to generate and supply energy by running the glycolytic pathway [10]. Adult schistosomes import exogenous glucose, equivalent to their dry body weight every 4 hours from host blood by using glucose transporters in their tegumental membranes [11,12]. In *C. sinensis*, glucose transporter and Na⁺/glucose co-transporter are expressed abundantly in the adult stage but less so in the metacercaria stage as presented in the *C. sinensis* transcriptome [13]. Adult *C. sinensis* worms uptake glucose to produce energy in the anaerobic environment of the bile duct [14]. Therefore, we expected that *C. sinensis* could be labeled with 2-deoxy-2-[¹⁸F]fluoro-D-glucose (¹⁸F-FDG), a glucose analogue used for the radiolabeling and diagnostic imaging of cancer cells [15]. Thus, by *ex vivo* labeling *G. neophila* with ¹⁸F-FDG, we hoped their migration in the final host could be traced *in vivo* by positron emission tomography-computed tomography (PET-CT).

In *vivo* imaging techniques have strong merits for the non-invasive tracing on pathogens moving within tissues of living animals, as they involve minimal manipulation and/or euthanasia of animals, and allow repetitive tracking in same animals. Furthermore, as was found in the present study, these techniques make it possible to monitor the distribution and migration of *C. neophila* *in vivo* from the duodenum to the liver or distal bowel. This study was carried out to determine how *C. neophila* find their way and how rapidly they migrate to the intrahepatic bile duct by using *in vivo* ¹⁸F-FDG radiolabeling and PET-CT in a rabbit model.

**Materials and Methods**

1. **Collection of *C. sinensis* metacercariae**

   Topmouth gudgeons (*Pseudorasbora parva*), the second intermediate host of *C. sinensis*, were purchased at a fish market in Shenyang, Liaoning Province, People’s Republic of China. Fish were ground then digested in artificial gastric juice (8 g of pepsin 1:10,000 (MP Biochemicals Co., Solon, OH, USA) and 8 ml of concentrated HCl in 1 liter of water) for 2 hr at 37°C [10]. To remove particulate matters, the digested soup was filtered through a sieve of 212 μm mesh. *C. sinensis* metacercariae (135–145 μm x 90–100 μm) were then filtered out using sieves of 106 and 53 μm meshes and washed thoroughly several times with 0.85% saline. *C. sinensis* metacercariae were collected under a dissecting microscope and stored in phosphate-buffered saline at 4°C until required [10].

2. **Labeling *C. neophila* with radio-isotope**

   The metacercarial cyst wall of *C. sinensis* is thick and can hinder glucose diffusion. Thus to maximize radiolabeling efficiency, metacercariae were excysted and juvenile worms were liberated from cysts. The *C. sinensis* metacercariae were excysted by treating them with 0.05% trypsin at 37°C for 5 minutes (Gibco, Grand Island, NY, USA) in 1× Locke’s solution (150 mM NaCl, 5 mM KCl, 1.8 mM CaCl₂, 1.9 mM NaHCO₃), a maintaining medium of *G. neophila* [16]. *G. neophila* were washed 5 times with 1× Locke’s solution, and used immediately. *G. neophila* were divided into two groups of 10–270 juveniles each; one was of *G. neophila* that excysted just before radiolabeling and the other was of the *G. neophila* fasted for 24 hours. The two *G. neophila* groups were radio-labeled with ¹⁸F-FDG by incubating them in 1× Locke’s solution containing 74 MBq ¹⁸F-FDG at 37°C for 15, 30, or 60 min. After washing 3 times with 1× Locke’s solution, radioactivity was measured for 1 min using a PET (GEMINI TF, Philips Healthcare, Cleveland, OH, USA). Numbers of *C. neophila* were counted and labeling efficiency was calculated as counts per minute (cpm) divided by number of the *C. neophila*. Radio-labeling efficiencies of the *C. neophila* in both groups were measured 3 times and significant differences were determined using the student’s *t*-test.

3. **Galbladder contraction in response to cholecystokin by cholescintigraphy**

   Rabbits (New Zealand White, male, 2.2–2.5 kg) were purchased from Samtako Bio Korea Inc. (Osan, Korea). Rabbits were cared for and handled according to guidelines issued by Chung-Ang University College of Medicine Animal Facility (an accredited facility) in accordance with AAALAC International Animal Care policy. Animal experiments were approved by the institutional review board of the Chung-Ang University animal facility (CAUMID 09-0024).

   Galbladder contraction and emptying time induced by cholecystokin-8 (CGK-8) varied from rabbit to rabbit. To select rabbits that responded sensitively to CGK-8, cholescintigraphy and ⁹⁹mTc-mebrofenin (3-bromo-2,4,6-trimethylphenyl carbamoylmethyl iminodiacetic acid) were used. Briefly, rabbits were fasted for 12 hrs and anesthetized with a 0.47 mg/kg Rompun (xylazine hydrochloride; Bayer Korea, Seoul, Korea) and 12.5 mg/kg Zoletil 50 (Zolazepam and Tiletamine; Virvac Korea, Seoul) using a 256×256-pixel matrix at an energy range of 20% at 140 keV.

4. **In vivo imaging of migration of the *C. neophila* using PET-CT**

   Fresh *G. neophila* (n = 3,000) were radio-labeled with ¹⁸F-FDG by incubating them in a maintaining medium containing 74 MBq ¹⁸F-FDG at 37°C for 15 min. *G. neophila* were washed 3 times with 1×...
Locke’s solution and then placed in 500 µl of 1 × Locke’s solution. The procedure was conducted as follows (Figure 1). A rabbit sensitive to CCK-8 was anesthetized with 0.47 mg/kg Rompun and 12.5 mg/kg Zoleit 50 by intramuscular injection and placed in restraints in a supine position on a plastic board. A catheter (3F Simmons II, Cook Co., Bloomington, IN, USA), equipped with a guidewire (0.035” Radilocus®, Terumo, Tokyo), was inserted through the animal’s mouth and its end positioned in the mid duodenum under guidance (Axiom Artis; Siemens, Erlangen, Germany). The rabbit was then moved with the catheter in situ and placed in PET-CT bed. To stimulate gallbladder contraction and bile juice release, 20 ng/kg of CCK–8 was injected intravenously every minute over this experiment [17]. After 12 minutes of CCK-8 injection, 18F-FDG-labeled CNEJs in 500 µl of 1 × Locke’s solution were introduced into the mid duodenum through the catheter; residual CNEJs in the catheter were flushed into the duodenum with 0.5 ml of 1 × Locke’s solution. One transmission CT image was obtained before the introduction of the 18F-FDG-labeled CNEJs and a dynamic PET scan then was performed over 90 min with a 3-min acquisition per frame. Finally, one static PET image was scanned for 10 min. This procedure is depicted schematically as a flow-chart in Figure 1.

All photon data were collected using a dedicated PET-CT scanner. PET images were reconstructed after applying CT-based attenuation and scattering corrections using the ordered subset expectation maximization algorithm (2 iterations, 16 subsets) for the presence of focal 18F-FDG uptake by radiolabeled CNEJs. Migration of the CNEJs to the intrahepatic bile ducts was determined using 99mTc-mebrofenin and cholescintigraphy. After 99mTc-mebrofenin injection, radioactivity increased immediately in the gallbladder to reach a peak at about 15 min, which was maintained for over 60 min (Figure 3A). When rabbits were injected intravenously with CCK-8, 99mTc-mebrofenin was rapidly released from the gallbladder and flowed down the small intestine (Figure 3B). Of the 16 rabbits tested for gallbladder contraction, 6 responded sensitively to CCK-8. On average, it took 11.5 min to evacuate 50% of the gallbladder volume after the first CCK-8 injection. The rabbits responding to CCK-8 were allowed one week to recover and were then included in the in vivo imaging experiments.

To confirm migration of the CNEJs to the bile duct, 10,760, 7,726 and 13,842 cpm/worm when incubated for 15, 30, and 60 min, and fasted CNEJs were labeled with 11,115, 8,043, and 12,318 cpm/worm when incubated for 15, 30, and 60 min, respectively (Figures 2A & B). Labeling efficiencies were similar in the two groups at all time points. For downstream experiments, fresh CNEJs were radiolabeled with 18F-FDG at 37°C for 15 min.

2. Gallbladder contraction and bile release

To determine an appropriate time point to inoculate the 18F-FDG-labeled CNEJs in the duodenum after CCK-8 injection, gallbladder contraction and 50% bile emptying times were determined using 99mTc-mebrofenin and cholescintigraphy. After 99mTc-mebrofenin injection, radioactivity increased immediately in the gallbladder to reach a peak at about 15 min, which was maintained for over 60 min (Figure 3A). When rabbits were injected intravenously with CCK-8, 99mTc-mebrofenin was rapidly released from the gallbladder and flowed down the small intestine (Figure 3B). Of the 16 rabbits tested for gallbladder contraction, 6 responded sensitively to CCK-8. On average, it took 11.5 min to evacuate 50% of the gallbladder volume after the first CCK-8 injection. The rabbits responding to CCK-8 were allowed one week to recover and were then included in the in vivo imaging experiments.

3. Monitoring of CNEJ migration

Under x-ray visualization and anesthesia, the end of a catheter was positioned in the mid duodenum (Figure S1). The rabbit was then anesthetized in the PET-CT bed; anesthesia was maintained with intravenous CCK-8 at a dose of 20 ng/kg every minute during PET-CT scanning. One abdominal CT image was obtained initially and then dynamic PET scanning was started. Three minutes after the initial PET scanning, the 18F-FDG-labeled CNEJs were inoculated into the mid duodenum (Figure 1). Dynamic and static PET scans were carried out using PET-CT on migrating 18F-FDG-labeled CNEJs in 6 rabbits, which included 2 controls.

Signals emitted from the 18F-FDG-labeled CNEJs were detected in the intestine of the 4 experimental rabbits by PET, and thus, we were able to trace CNEJ migration by in vivo imaging. When the 18F-FDG-labeled CNEJs were injected through the catheter, signals were detected at end of the catheter in the duodenum and along the small intestine driven by peristalsis.

Results

1. Labeling efficiency of CNEJs with 18F-FDG

Fresh CNEJs were labeled with 10,760, 7,726 and 13,842 cpm/worm after incubation in radiolabeling media for 15, 30, and 60 min, and fasted CNEJs were labeled with 11,115, 8,043, and 12,318 cpm/worm when incubated for 15, 30, and 60 min, respectively (Figures 2A & B). Labeling efficiencies were similar in the two groups at all time points. For downstream experiments, fresh CNEJs were radiolabeled with 18F-FDG at 37°C for 15 min.

Figure 1. Flow-chart of the PET-CT imaging of the migration of newly excysted Clonorchis sinensis juveniles (CNEJs) in rabbits.
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along the distal portion of the intestine (Figure S2). Signals of CsNEJs appeared in the liver as early as 7–9 min after inoculating the 18F-FDG-labeled CsNEJs into the duodenum (Figure 4A). As time elapsed, some photon spots emerged in the liver region and enlarged whereas others faded. These spots appeared to be randomly and evenly distributed in the liver regardless of lobe structure (Figures 4A–D), and gradually increased in number to plateau at about 21 min after inoculation of the radiolabeled CsNEJs (Figures 4 & 5). Spots suggestive of CsNEJs moving through the common bile duct were not observed in PET-CT images. In static PET-CT images taken finally over 10 min, CsNEJs appeared to aggregate in central region of the liver (Figures 4E & F). Of the CsNEJs inoculated into the duodenum, some migrated up to the bile ducts and others down to the lower bowel driven by peristalsis (Figure S2).

In rabbits not injected with CCK-8 (the negative control group), signals of 18F-FDG-labeled CsNEJs were only observed in the small intestine in dynamic and static PET images.

At 4 weeks after the CsNEJs inoculation into the duodenum, adult C. sinensis worms were found to inhabit and to have provoked pathologic changes in the bile ducts. On average 1,077±806 adults were recovered from the biliary tracts of the rabbits (Figure S3).

**Discussion**

*In vivo* the migration route of *C. sinensis* was indirectly determined by ligating the common bile ducts of hosts. Recently, live *Schistosoma mansoni* adults in mice were labeled with protease-activated fluorochrome or 18F-FDG and visualized, localized, and quantified using fluorescence molecular tomography or PET [18,19]. In the present study, we applied the methodologies and investigated PET-CT as a new *in vivo* imaging method for monitoring the migration of CsNEJs and their localization in the rabbit liver. The rabbits are highly susceptible to and retain the *C. sinensis* infections long time to evaluate impact of the infection on the hepatobiliary system. The rabbits have the biliary system similar to that of human. Distribution of *C. sinensis* in the liver of the experimental rabbits was proportional to volume of the liver lobes [20–22]. We, therefore, expected the rabbit as a reliable experimental animal model to study bile-chemotactic migration of...
the CsNEJs, suggesting that findings obtained from the rabbits are applicable to human.

Trematodes import glucose through glucose transporter, and a large number of glucose transporters have registered in the *C. sinensis* transcriptome database [13]. $^{18}$F-FDG is a glucose analog tagged with isotope $^{18}$F, and is transported into cytoplasm by glucose transporters in cell membrane. In the cytoplasm, FDG is phosphorylated to FDG-6-phosphate by hexokinase, and FDG-6-phosphate is neither metabolized further nor able to diffuse out of cells. Thus, FDG-6-phosphate is trapped and accumulates in cells as the dephosphorylation of FDG-6-phosphate by glucose-6-phosphatase in cytoplasm is a slow process [15,23]. We expected that fasted CsNEJs would uptake more FDG than fresh CsNEJs because CsNEJs should have consumed their reserve energy source, primarily glucose, during fasting in glucose-free 1× Locke’s solution. However, FDG uptakes in both groups were similar, suggesting that FDG moved

**Figure 4. PET-CT images of photons emitted from $^{18}$F-FDG-labeled CsNEJs in the rabbit liver.** Photon images represent *Clonorchis sinensis* arriving at the intrahepatic bile ducts of the rabbit liver. A–D, dynamic images at 7–9, 16–18, 22–24, and 52–54 minutes after CsNEJs inoculation. Static axial (E) and coronal (F) images taken 90 minutes after the CsNEJs inoculation. The liver is located as a region of interest (circle).

**Figure 5. Plot of photon counts from the rabbit liver.** The $^{18}$F-FDG-labeled CsNEJs were attracted with bile juice released by CCK-8-induced gallbladder contraction.

![Figure 4](https://example.com/figure4.png)

![Figure 5](https://example.com/figure5.png)
quickly into the tegument of CsNEJs through glucose transporter by facilitative diffusion, as was observed for schistosomes [24,25].

During our studies, we have observed that CsNEJs move toward bile dose-dependently by chemotaxis in in vitro assays (unpublished data). Based on our data and the notion that C. sinensis juveniles migrate up through the common bile duct, it was essential that bile juice is released from the gall bladder to attract CsNEJs into the common bile duct.

Technetium labeled hepatobiliary radiopharmaceuticals has greatly facilitated studies on gallbladder function [26]. Since CCK-stimulated cholecintigraphy was first reported in 1979, gallbladder emptying function has been measured by using standard chologagogic stimulus agents by biliary excretion scintigraphy [27,28]. Cholescintigraphy with 99mTc-9mTc-iminodiacetic acid has been used to diagnose diseases in the biliary system, such as, bile duct obstruction, cholelithiasis, cholecystitis, and biliary fistula [29–31].

The gallbladder normally fills with hepatic bile during fasting and empties its contents into the duodenum in response to stimulation by CCK, either released endogenously following a meal or administered exogenously [32]. However, gallbladder emptying response to exogenous CCK varies among patients and experimental animals. In this study, gallbladder contraction and bile juice release was achieved by repeatedly injecting CCK-8. By cholecintigraphy, 99mTc-mebrofenin was found to be released rapidly from gallbladders after CCK-8 administration. Thus, this scheme enabled us to study in vivo bile-chemotactic behavior of CsNEJs in rabbits.

Using CsNEJ radiolabeling and bile excretion from gallbladder, images of CsNEJs migrating to the intrahepatic bile ducts in rabbits were obtained by PET-CT. The radionabeled CsNEJs were inoculated into the mid duodenum, which is supposed to be an excystation site for C. sinensis metacercariae [3,6]. We visualized 18F-FDG-labeled CsNEJs migrating to the liver in experimental rabbits using PET-CT. The first signals of CsNEJs arriving at the liver from the duodenum were detected by dynamic PET as early as 7–9 min after inoculating CsNEJs into duodena. At 21 minutes post-inoculation, photon signals emitted from CsNEJs in liver appeared to have stabilized though their intensities undulated, which suggested most CsNEJs responsive to bile immediately migrated up to the intrahepatic bile duct. Imaging was ended with a final static PET-CT image because signals were of greater intensity than on dynamic PET images, suggesting that some CsNEJs were late to arrive and accumulated in the intrahepatic bile ducts [3]. In in vitro assays, CsNEJs showing rapid bile-taxis were promptly re-activated and moved rapidly and continuously toward bile added to assay chambers, and slow responders responded slowly (unpublished data).

The artificial manipulation of CsNEJs employed in this study, that is, in vitro excystation and radiolabeling, and inoculation into the duodenum, may have reduced adaptation to body temperature, chemotactic response to bile, migration to the bile duct, and survival in bile juice. To compensate for this, in the present study, 3–5 times more 18F-FDG-CsNEJs than normally usual experiment was inoculated via catheter into the duodenum. We believe that slow responders arrived late at the intrahepatic bile ducts after PET scans, and increased numbers of C. sinensis adult worms recovered from the bile ducts [3]. When filet of the fresh water fish was minced by teeth and ingested by mammalian animals including human, the C. sinensis metacercariae could be released from the filet in the stomach after 1–2 hour, and then passed down to the duodenum. Considering immediate excystation of the C. sinensis metacercariae in contact with trypsin [6], human infection may take place within 2–3 hours after eating raw filet of the fresh water fish.

We searched for phonic signals from the common bile ducts in dynamic and static PET-CT images of experimental rabbits, but found no signal. The common bile duct is narrow and CsNEJs either passed rapidly or steadily in file, and thus, only small number of juveniles (not enough to create a PET-CT image) was captured in a given frame. Furthermore, anatomically the common bile duct is located in the deep abdomen under the liver, which hinders emitted photons.

Collectively, we report for the first time that CsNEJs were efficiently radiolabeled in vitro with 18F-FDG, and that CsNEJs migrate quickly with bile-chemotaxis to the intrahepatic bile duct as visualized in rabbits by PET-CT.

Supporting Information

**Figure S1** Insertion of a catheter into the mid duodenum of a rabbit under anesthesia. (TIF)

**Figure S2** PET-CT coronal images showing some 18F-FDG-labeled CsNEJs driven down the small intestine by peristalsis. A-D, 9, 18, 24, and 27 minutes after inoculating radiolabeled CsNEJs into the mid duodenum. (TIF)

**Figure S3** Clonorchis sinensis from an experimental rabbit liver 4 weeks after a bile-chemotaxis experiment. A, Adult flukes in the rabbit liver, hematoxylin-eosin stained. B, Adult flukes recovered from the rabbit’s liver. (TIF)

Author Contributions

Conceived and designed the experiments: S-JH. Performed the experiments: TIK WGY BKK J-WS. Analyzed the data: TIK BKK J-WS S-JH. Contributed reagents/materials/analysis tools: BKK J-WS S-JH. Wrote the paper: TIK BKK J-WS S-JH.

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