An Integrated Quad-Modality Molecular Imaging System for Small Animals

Yanye Lu1, Kun Yang2, Kedi Zhou1, Bo Pang1, Guohe Wang1, Yichen Ding1, Qiushi Zhang1, Hongbin Han1, Jiahe Tian3, Changhui Li1, and Qiushi Ren1

1Department of Biomedical Engineering, College of Engineering, Peking University, Beijing, China; 2Department of Control Technology and Instrumentation, College of Quality and Technical Supervision, Hebei University, Baoding, China; and 3Department of Nuclear Medicine, The Chinese PLA General Hospital, Beijing, China

We developed a novel integrated quad-modality system that included 3 molecular imaging methods (PET, SPECT, and fluorescence molecular imaging [FMI]) and 1 anatomic imaging modality (CT). This system could study various biologic processes in the same animal using multiple molecular tracers. In addition to the technology development, we also discussed the optimization strategy of the imaging protocols. The performance of this system was tested, and the in vivo animal experiment showed its power to trace 3 different molecular probes in living tissues. Our results demonstrated that this system has a great potential for the preclinical study of diseases. Methods: A prototype system integrating PET, SPECT, CT, and a charge-coupled device-based free-space FMI system has been developed. Imaging and fusion capabilities of the system were evaluated by a multimodality phantom. In addition, a mouse disease model with both tumor and inflammation was studied by this system to examine the in vivo performance. The 3 types of molecular probes—18F-FDG, 99mTc(HYNIC-3PRGD2)(tricine)(TPPTS) (99mTc-3PRGD2) (HYNIC = 6-hydrazinonicotinyl; TPPTS = trisodium triphenylphosphine-3,3′,3″-trisulfonate; 3PRGD2 = PEG3-E[PEG3-c (RGD4K)], and 3-(triethylxosylyl) propyl-Cy7-entrapped core-cross-linked polymeric micelle (Cy7-entrapped CPM) nanoparticles—were used to target 3 different biologic processes in the tumor caused by pulmonary adenocarcinoma A549 cells. Moreover, the strategy to optimize multimodal molecular imaging procedure was studied as well, which could significantly reduce the total imaging time.

Results: The imaging performance has been validated by both phantom and in vivo animal experiments. With this system and optimized imaging protocol, we successfully differentiated diseases that cannot be distinguished by a single molecular imaging modality. Conclusion: We developed a novel quad-modality molecular imaging system that integrated PET, SPECT, FMI, and CT imaging methods to obtain whole-body multimodality images of small animals. The imaging results demonstrated that this system provides more comprehensive information for preclinical biomedical research. With optimized imaging protocols, as well as novel molecular tracers, this quad-modality system can help in the study of the physiology mechanism at an unprecedented level.

Key Words: multi-modality imaging; animal imaging; molecular imaging; instrumentation

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Molecular imaging traces the biologic processes at the molecular level in living tissues with high resolution and specificity, sensing diseases at their very early stages. Since it was proposed in the 1990s, molecular imaging has become a key tool in biomedical research and clinical imaging (1,2). Various molecular imaging methods based on different mechanisms, such as radio-labelling and optical labeling, have gained significant progress over the past decades (2–11).

The biologic system involves complex biochemical reactions and the transfer of various cells and molecules. For instance, a breast cancerous lesion could contain several subtype cancer cells, which behave differently in metastasis, metabolism, and angiogenesis. Information on the development and transportation of these cells is essential for accurate diagnosis and effective treatment. In addition, different biologic processes could have serious overlap in molecular reactions. One example is that both cancer and inflammation can cause abnormal metabolic activities, so that tracing glucose metabolism alone is not suitable to separate these 2 distinct diseases. Thus, multiple probes targeting different biomarkers are desired to get important complementary information about these multiple biologic processes. Because different kinds of molecular probes rely on different detecting mechanisms, the integrated multimodal molecular imaging system that can trace multiprobes gains increasing interest and will have great impact for biomedical imaging (7,12).

Up to now, 3 major molecular imaging modalities—PET, SPECT, and fluorescence molecular imaging (FMI)—have been widely used. Relying on radioactive tracers, PET and SPECT have already gained significant clinical applications (12–16). Fluorescence imaging requires optical tracers (e.g., green fluorescence protein) and is favored by researchers because of its versatility, low-cost, and ease of manufacture and use and more importantly because of its unique genetically coding capability. Furthermore, when the image coregistration, safety, reliable handling, and minimization of total imaging time are considered, fluorescence imaging is desired for multimodal molecular imaging in a single system. There are a few existing multimodality animal imaging systems (15) as well as commercial trimodality systems, such as the Inveon (Siemens) (14), FLEX Triumph (Gamma Medica-Ideas), and Albira (Carestream), that have...
MATERIALS AND METHODS

Quad-Modality Molecular Imaging System

The quad-modality imaging system used a modular design—that is, PET, CT, SPECT, and FMI modalities were designed and developed separately (more details are described in the supplemental data). We used a line-type x-ray detector with a helical scan mode in the CT module and a rotary transmission module design. The SPECT system, which has 2 parallel-opposed γ cameras to increase the axial field of view that can cover the whole-body area of most mice, was mounted on the same rotating plate with CT (Fig. 1). The γ camera had a 22 × 22 cm2 cerium-doped lutetium-yttrium oxyorthosilicate scintillator array coupled to a Hamamatsu H8500 position-sensitive multianode photomultiplier tube by optical silicone oil. The PET individual module (20) was a fixed ring containing 54 silicon photomultiplier-based detectors. On the end side of the system, a noncontact, full-angle and vertically rotating FMI module was installed. The FMI module can perform not only traditional fluorescence 2-dimensional imaging with both epillumination and transillumination but also tomographic 3-dimensional imaging. In addition, a transparent animal bed was mounted on a high-precision motor-controlled 1-dimensional translation stage, which carried the animal from one modality to another automatically. The PET, SPECT, and CT modules were set up on one side of the motor-controlled animal bed, and the FMI module was placed on the other end because that module required strictly light shielding. The entire imaging system can be installed on a 900 × 1,500 mm optical table (Fig. 1B). There was a custom-designed enclosure (not shown in the figure) that was used to cover the entire system and that also optically shielded the FMI subsystem. In this way, this design not only achieved a compact multimodality system but also avoided signal conflict and optical shielding. Figure 2 presents the overall system design, which consisted of rotary transmission, electronics, power supplies, and several pieces of supporting equipment. The animal bed can be shifted between different modalities, and the multimodality imaging fusion can be done precisely with the coregistered animal position. Geometric calibration and image coregistration were processed using fiducial markers on the animal holder.

Optimizing Multimodality Imaging Protocols

Both phantom and animal experiments were performed to evaluate the imaging capability and fusion performance of the system. Because there are vast differences in the uptake period and imaging technique, each molecular imaging modality requires a specific procedure. Therefore, the overall imaging time can last up to 6 h in our system without optimization of the imaging protocols. However, long imaging time not only brings difficulties in dynamic study, but also may cause a health hazard for the animals. Thus, optimization of the imaging protocol for multimodal molecular imaging is greatly desired. Given a physiologic or pathologic study case, the following are several primary issues to be considered in assigning the imaging sequence:
with 7.4 MBq (0.2 mCi) of $^{18}$F-FDG and PET scanning was performed (180° rotation, scanned for 10 s/3°, 60 angles in total); afterward, the animal bed was shifted to the field of view for the PET/SPECT/CT modules and subjected to SPECT scan (360° rotation, scanned for 4 s/10°, 36 angles in total). There were 2 holes into which glass tubes (inner diameter, 2.2 mm; length, 50 mm) with radiolabeling medicines and fluorescence dyes could be inserted at different insertion depths (28 and 45 mm) in this cylinder. The bottom zones of the glass tubes were injected with 20 μL (10 μM) of indocyanine green as the fluorophore for FMI; the middle section of the 45-mm-insertion-depth glass tube was injected with 7.4 MBq (0.2 mCi) of $^{99m}$Tc-2-methoxyisobutylisonitrile. The phantom was placed on the animal bed to be first imaged by fluorescence molecular tomography (360° rotation, scanned for 4 s/10°, 36 angles in total); afterward, the animal bed was shifted to the field of view for the PET/SPECT/CT modules and subjected to SPECT scanning (180° rotation, scanned for 10 s/3°, 60 angles in total). Then, the middle section of the 28-mm-insertion-depth glass tube was injected with 7.4 MBq (0.2 mCi) of $^{18}$F-FDG and PET scanning was performed (300- to 650-keV energy window and 12-ns timing window, scanned for 5 min/per bed position, 3 bed positions). Finally, the phantom was moved to the field of view for the CT scan (60 slices, 1.32-mm helical pitch, 360 projection numbers, and 25-ms sampling intervals; x-ray tube voltage, 50 keV, at 10 mA).

FMI data were reconstructed using the algebraic reconstruction technique with a relaxation parameter of 0.1 and 200 iterations. Both SPECT and PET data were reconstructed using the 2-dimensional ordered-subsets expectation maximization algorithms with 10 iterations and 16 subsets; CT data were reconstructed using the filtered back-projection algorithm with a Butterworth filter of order 3 and a cutoff at 0.1 of the Nyquist frequency. All the reconstructed images were coregistered through the geometric calibration to perform image fusion.

**Phantom Studies**

The evaluation of imaging and coregistration performance for the prototype system was done by a custom-made multimodality phantom study. The phantom was a solid cylinder (diameter, 30 mm; length, 57 mm) having a light scattering property similar to tissue (Fig. 3A). There were 2 holes into which glass tubes (inner diameter, 2.2 mm; length, 50 mm) with radiolabeling medicines and fluorescence dyes could be inserted at different insertion depths (28 and 45 mm) in this cylinder. The bottom zones of the glass tubes were injected with 20 μL (10 μM) of indocyanine green as the fluorophore for FMI; the middle section of the 45-mm-insertion-depth glass tube was injected with 7.4 MBq (0.2 mCi) of $^{99m}$Tc-2-methoxyisobutylisonitrile. The phantom was placed on the animal bed to be first imaged by fluorescence molecular tomography (360° rotation, scanned for 4 s/10°, 36 angles in total); afterward, the animal bed was shifted to the field of view for the PET/SPECT/CT modules and subjected to SPECT scanning (180° rotation, scanned for 10 s/3°, 60 angles in total). Then, the middle section of the 28-mm-insertion-depth glass tube was injected with 7.4 MBq (0.2 mCi) of $^{18}$F-FDG and PET scanning was performed (300- to 650-keV energy window and 12-ns timing window, scanned for 5 min/per bed position, 3 bed positions). Finally, the phantom was moved to the field of view for the CT scan (60 slices, 1.32-mm helical pitch, 360 projection numbers, and 25-ms sampling intervals; x-ray tube voltage, 50 keV, at 10 mA).

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**In Vivo Animal Studies**

Besides the phantom study, the mouse disease model was studied using this quad-modality imaging system. We induced a xenograft tumor (induced by hypodermic injection of human pulmonary adenocarcinoma A549 cells [$2.5 \times 10^7$, 2.5 mL]) and chronic inflammation (induced by hypodermic injection of the proinflammatory substance of Bacillus Calmette-Gu in the right shoulder [1 mg/mL] and the right hindlimb [0.2 mL]) of a 16-wk-old, 20-g BALB/C nude mouse. We used $^{18}$F-FDG for PET imaging, $^{99m}$Tc-3PRG$_2$ (21) for SPECT imaging, and Cy7-entrapped CCPM (22,23) nanoparticles for FMI. $^{18}$F-FDG traces high-glucose-using cells, $^{99m}$Tc-3PRG$_2$ is able to target integrin αvβ3-positive tumors, and Cy7-entrapped CCPM nanoparticles can preferentially accumulate in the tumor site because of the enhanced permeability and retention effect. The mouse was intravenously injected with 0.2 mL (1.5 $\times 10^{13}$ particles/mouse) of Cy7-entrapped CCPM nanoparticles and 37.0 MBq (1.0 mCi) of $^{99m}$Tc-3PRG$_2$ via the tail vein. After a 50-min uptake period, the mouse underwent anesthesia (2.50% isoflurane in oxygen [1.5 l/min] for inducing anesthesia and then down to 1.00% isoflurane in oxygen [0.8 l/min] for maintenance). The mouse was scanned with SPECT (180° of rotation, 20 s/3°, 60 angles; duration, 20 min) after the 60-min uptake period of $^{99m}$Tc-3PRG$_2$. Then the mouse was intravenously injected with 37.0 MBq (1.0 mCi) of $^{18}$F-FDG via the tail vein, and the anesthesia gas was increased to 2.5% isoflurane in oxygen during the injection and down to 1.00% after the injection. At the end of the 120-min uptake period of Cy7-CCPM nanoparticles, a helical CT scan (60 slices; 1.5-m helical pitch; 360 projection numbers and 25-ms sampling intervals; x-ray tube voltage, 50 keV; tube current, 10 mA) was obtained. Then, the mouse underwent FMI (epiillumination mode; integration times, 5 s). Both CT imaging and FMI were performed during the 60-min $^{18}$F-FDG uptake period. After the period, the mouse was PET-scanned (300- to 650-keV energy window, 12-ns timing window, 10 min/bed position) for 40 min of 4 bed positions. The whole animal experimental procedure is illustrated in Figure 4B. After the examination, the mouse was sacrificed, and the tumor and inflammation tissues were excised, fixed in 4% paraformaldehyde, and embedded in paraffin. Hematoxylin and eosin staining was performed on 5-μm-thick tissue sections.

The PET/SPECT/CT images were reconstructed as aforementioned. Because the subcutaneous tumor was located in a shallow region of the body, only the 2-dimensional fluorescence imaging was given by using the epiillumination FMI. All the reconstructed images were coregistered automatically to perform image fusion. Animal studies were performed in accordance with the guidelines from the Peking University Laboratory Animal Centre and protocols approved by this institution.

**RESULTS**

Performance Evaluation by Phantom Study

Figure 3 shows the imaging results, including images by individual modalities and coregistered images. Both PET and SPECT provided high-quality images without observable crosstalk artifacts. In order to demonstrate the spatial resolution, a length of bubble (3 mm) separating 2 PET medicine drops was made, and this small gap was clearly distinguished in PET images. In addition, FMI also successfully reconstructed
the fluorophores deep in the phantom. In Figure 3C, the maximum-intensity-projection rendering of the fused images demonstrated the accurate coregistration. The imaging results indicated that our hybrid multimodality imaging system could provide appropriate imaging and accurate image fusing.

**In Vivo Animal Studies**

Figure 5 shows the multimodality and multiprobe imaging results. Complementing the CT results, the PET images showed high radioactivity accumulation in the brain, right shoulder, and right legs. In SPECT images, the radioactivity accumulation of 99mTc-3PRGD2 primarily occurred in the right shoulder, abdomen, and area of bladder. In addition, according to the FMI results in Figure 4B, only the area of the right shoulder had a significant accumulation of Cy7-entrapped CCPM nanoparticles. From either the PET image or the SPECT image alone, the tumor area remained unclear. However, we were able to identify the tumor’s location by the complementary information obtained from the PET–SPECT–CT fusion images. On the other hand, fluorescence molecular images showed the correct area that was suggestive of tumor, but the imaging resolution was low. Complementing FMI with radioactive molecular imaging significantly improved both specificity and resolution. This result successfully demonstrated the superior power of multimodal molecular imaging.

Moreover, by following the optimized imaging procedure, the entire imaging time for this in vivo study, including preparation and imaging, took only 3 h. However, the procedure can last up to 6 h just to sum all imaging times from each individual modality. The significant reduction in imaging time is essential to study dynamic physiologies. The imaging time can be reduced even further after system improvement.

**DISCUSSION**

Successful multimodal molecular imaging requires innovations in both technology development and imaging protocols. In this study, we not only developed the first quad-modality imaging system for preclinical in vivo molecular imaging studies, but also provided a strategy to optimize the imaging protocol. The performance of the system and image coregistration have been validated by both phantom and in vivo animal experiments.

In the animal disease model study, the quad-modality imaging system successfully found tumors and acquired images non-

![FIGURE 4. Timeline schematic of whole animal experimental procedure. (A) Typical quad-modality imaging procedure in this animal study takes more than 320 min, considering time needed for drug metabolism. (B) After optimization of imaging protocol, total time was reduced to 180 min, as well as avoiding crosstalk issue.](image-url)
invasively and in vivo. The 4 modalities of PET, SPECT, FMI, and CT can provide more comprehensive information for molecular imaging studies, improving the overall specificity. In our study, the image fusion of a tumor xenograft mouse (Fig. 5) showed complementary information that helped to differentiate between the tumor location and the inflammation location. These findings were confirmed by ex vivo histology.

This quad-modality system also has a great potential for cancer subtype diagnosis. Metabolism, metastasis, and neovascularization are the 3 most important characteristics of the pathologic behavior of cancer cells. According to recent studies, many cancers have various subtypes, and not all of them are aggressive. Differentiation of the subtypes of cancer cells has an important role in the study of the pathology of cancer as well as its treatment (24–26). Traditionally, molecular biology approaches such as immunohistochemistry are used for revealing the biologic behavior of cancer cells; however, recent molecular imaging approaches—which allow for noninvasive in vivo imaging and quantification of biologic processes such as metabolism, metastasis, and neovascularization—are more and more widely used in these studies. With the quad-modality system, 3 different tracers for PET, SPECT, and FMI can trace these 3 types of cells with high specificity and accurate image fusion. However, this cannot be done with any current single system (15,18).

In addition, this quad-modality imaging system was modular-designed. All 4 modalities were coaxial-aligned, and a transparent animal holder controlled by a high-precision translational and rotational stage can deliver the animal from one modality to another. Our system can be used as single, dual-, tri-, or quad-modalities such as CT, PET/CT, SPECT/CT, FMI-CT, PET–SPECT/CT, or PET–SPECT–FMI–CT, based on different research requirements.

Multimodal molecular imaging generally required an extended time. For instance, in our system, SPECT typically needs to wait for 60 min after the injection of medicine, and it took another 20–30 min to finish the scanning. In addition, different imaging modalities might have conflicts, such as the potential risk of crosstalk between PET and SPECT. However, summation of operation time by all 3 molecular imaging modalities, as well as the uptake time of radioactive medicine, will be about 6 h. Thus, the procedure of multimodal molecular imaging must be carefully optimized to minimize the total scanning time. To avoid crosstalk, the PET tracer was injected after the SPECT scan in our system. In addition, long-lasting fluorescence tracers could be injected the day before the experiment. Finally, we achieved a 3-h imaging time. The total length of the scanning time can be even shortened by performance-improvement of the imaging system. For example, we can inject both radiolabeling tracers simultaneously using a better SPECT detector that can differentiate signals from PET tracer (16). In addition to the technical development, the in vivo animal study is our next primary research work.

CONCLUSION

We developed a novel quad-modality molecular imaging system that integrated PET, SPECT, FMI, and CT imaging modalities to obtain whole-body multimodality molecular images of small animals. The optimization in both system setup and imaging procedure are all carefully considered. We demonstrated that this system provides more comprehensive information for preclinical biomedical research. Using optimized imaging protocols and novel molecular tracers, this quad-modality system can help people understand the physiology mechanism on an unprecedented level.

DISCLOSURE

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