Cell tracking using multimodal imaging

Mangala Srinivasa, Ignacio Melerob, Eckhart Kaempgenec, Carl G. Figdoranda and I. Jolanda M. de Vriesa*  

1. INTRODUCTION  
Multimodal imaging is the use of more than one imaging modality for a specific purpose, in this case, cell tracking. Cell tracking consists of following specific cells in vivo; this is often in terms of their localization, but can also be in terms of their fate, functionality or differentiation. Cell tracking using imaging has several advantages, such as the noninvasive nature of imaging which can allow longitudinal follow-up of cells, and the ability to acquire specific information such as about the numbers of cells in a region of interest, their viability and their functionality. Noninvasive imaging is also applicable to humans. For all these reasons, imaging plays a key role in the optimization of cellular therapeutics (1). Multimodal imaging is particularly powerful in that it allows for comprehensive monitoring of labeled cells, as the strengths of the different imaging modalities can be maximized. Multimodal imaging can be done using a single label or tracer that is visible using different imaging modalities, or a combination of imaging labels. The imaging modalities typically applied to cell tracking include whole body scintigraphy, magnetic resonance imaging (MRI), positron emission tomography (PET), fluorescence imaging (FLI), bioluminescence (BLI) and single photo emission computed tomography (SPECT).  

SPECT, PET and scintigraphy require the use of radioactive labels. For example, SPECT and scintigraphy are commonly done using 111In and 99mTc. These have half-lives of up to 3 days, dependent on the isotope. PET tracers, often 18F analogs of glucose or thymidine, are much more short-lived, with half-lives in the order of hours. Furthermore, all these agents result in radiation exposure to subjects and thus their dosage and repeated use is limited and tightly regulated. However, nuclear medicine imaging techniques such as these can be very sensitive and yield quantitative information on label content. MRI, on the other hand, does not require the use of radiolabels. Instead, cells are labeled with iron oxide nano- or microparticles, or other metals such as gadolinium (Gd) or manganese (Mn). MR imaging works on the 1H nucleus, present abundantly in biological tissues, particularly in the form of mobile water, resulting in exquisite anatomic detail. Contrast agents result in the labeled cells presenting as hyperintense (T1 agents) or hypointense regions (T2 and T2* agents) on appropriately weighted MR images. Imaging using certain iron oxide agents can be very sensitive, even up to single cell imaging (2). Note that the use of MR contrast agents is also closely regulated, and they are not officially approved as cell labels. MR reporter genes have also been developed, where the genes are typically involved in intracellular iron levels (3). However, quantification of cell numbers with contrast agents can be difficult. Recently, much work has focused on labeling cells with 19F agents for cell tracking. 19F MRI is quantitative, but suffers from sensitivity issues and is still in preclinical testing (4). Finally, the light-based in vivo imaging modalities, FLI and BLI, are perhaps the most common owing to their relative ease of use. However, the techniques are limited by the penetration depth of light, and are thus most often used in preclinical models, particularly nude mice, with limited potential for translational application to humans. The use of fluorescent agents to label cells is very well established, for example in microscopy and flow cytometry. For these reasons, fluorescent agents are most often combined with imaging agents for MRI, SPECT or PET. BLI is similar to FLI, except that it requires the expression of an enzyme, luciferase, which catalyzes the
Biography

Mangala Srinivas did her B.Sc. (Honours) at the National University of Singapore. From there, she went on to complete her Ph.D. at Carnegie Mellon University in Pittsburgh, USA. She has since been working at the Dept. of Tumor Immunology at the Nijmegen Center for Molecular Life Sciences (The Netherlands). Mangala’s initial work during her Ph.D. involved the development and application of 19F magnetic resonance imaging (MRI) for in vivo cell tracking. Her present work involves the development and application of multimodal imaging agents to various applications, both clinical and preclinical. Mangala is currently funded by a personal VENI grant from the Netherlands Institute for Scientific Research (NWO). She has recently been awarded a Starting Grant by the European Research Council (ERC).

Biography

Ignacio Melero graduated as MD from the University of Navarra School of Medicine and was trained as a resident in clinical immunology at Hospital de la Princesa (Universidad Autonoma de Madrid). He also earned a PhD degree working with Dr. Miguel Lopez Botet pioneering the characterization of NK cell inhibitory receptors. In 1994 he moved to Seattle (USA) where he worked with Karl E. Helström and Lieping Chen in tumor immunology. His work was devoted to study T cell ignorance of tumor antigens and the role of costimulation in mouse models of cancer. His studies of that time on CD137-mediated stimulation of curative antitumor immune responses have received much attention. In 1998 he returned to Pamplona (University of Navarra), where he works at the Clinica Universitaria and at the investigation centre (CIMA). He was appointed full professor of Immunology in 2004. His current areas of research are focused on from bench to bedside translational research with cell, gene and monoclonal antibody-mediated therapies for cancer.

Biography

Dr. Kaempgen has been working in the field of Dendritic Cells since more than 20 years and continuously contributed to both, basic research on the immunobiology of DCs and translational research resulting in DC based treatment of tumor patients (>100 peer reviewed publications). He received his MD in 1987 at the medical school of the LMU Munich and, before starting his dermatology fellowship at the Univ. of Würzburg (Günter Burg, Eva Bröcker) in 1990, was trained as research associate at the Depts. of Immunology, Munich (Gert Riethmüller) and Dermatology, Univ. of Innsbruck (Gerold Schuler, Peter Fritsch). From the very first beginning Dr. Kaempgen was involved in the development of today’s standard method to generate human DCs from peripheral blood monocytes and especially strategies to load DC with tumor antigens by RNA transfection, which was developed in close cooperation with Eli Gilboa at the Duke University, Durham. Following a position as Director of Immunology, Merix Inc., Durham, NC Dr. Kaempgen in 2003 became Associate Professor for Strategies of Cellular Immunotherapy at the Dep. Dermatology, Univ. of Erlangen (head: Gerold Schuler), where he currently is the responsible senior dermatologist for skin tumor patients and heading the interdisciplinary melanoma tumor board. Starting September on 2013 Dr. Kaempgen will be working as a partner at the dermatologikum in Berlin (www.dermatologikum-berlin.de).

Biography

Carl Figdor (Radboud University Medical Centre Nijmegen) obtained his Masters from the University of Utrecht, and his PhD degree from the University of Amsterdam working at the Netherlands Cancer Institute, where he was tenured in 1985 and started his own research group on Tumor Immunology. In 1992 Carl Figdor became Professor in Cell Biophysics at the University of Twente, and in 1994 he moved to Nijmegen to start a new department on Tumor Immunology as a Professor in Immunology. From 2001 – 2011 he served as the first scientific director of the Nijmegen Centre for Molecular Life Sciences (NCMLS). Carl Figdor is a member of the Royal Netherlands Academy of Arts and Sciences and is the recipient of several awards, including the Dutch Spinoza Award in 2006, and the KWO award from the Dutch Cancer Society in 2009. He obtained an ERC Advanced Investigator grant in 2011.

Biography

Jolanda de Vries is a Professor at the Department of Tumor Immunology at the Nijmegen Centre for Molecular Life Sciences (NCMLS) at the Radboud University Medical Center. She was one of the pioneers in the translation of dendritic cell (DC) biology into potential clinical application, initiating the first clinical phase I/II studies in which patients were vaccinated with DCs loaded with tumor-specific peptides in 1997. Jolanda was also the first to apply MRI to clinical cell tracking in 2005. More recently, she has applied PET to study DC functionality in patients after transfer. Clinical trials involving DC vaccination and a combination of imaging modalities to effectively monitor the cells in patients are in process.
conversion of a substrate (luciferin to oxyluciferin) with the release of a photon. The photon released is detected. Typically, cells express luciferase as a reporter gene and luciferin is injected intravenously. Thus, this technique requires that the relevant cells be viable for imaging, not simply the presence of label. Reporter genes can also be used to label cells for PET imaging where a $^{18}$F-labeled precursor is metabolized and retained by the gene-expressing cells (5).

2. ROLE OF MULTIMODAL IMAGING

2.1. Validation

One of the main advantages of multimodal imaging is that an established imaging modality or technique can be combined with a more experimental one. Typically FLI and fluorescence-based techniques, such as microscopy, histology and flow cytometry, function as the established modality and can be considered the gold standard. Fluorescent dyes are well-characterized and relatively easy to add to imaging agents, for example, simply dissolved in a surfactant with liposomal agents or covalently bound to particulate agents. Alternatively, some studies use transgenic cells that express a fluorescent protein, or luciferase for BLI or a reporter enzyme for PET. These can be constitutionally expressed or used as a reporter gene for a given biofunction. In either case, luminescence indicates live and functional cells, and not simply the presence of imaging label, when the expressed luciferase is ATP-dependent. A recent example demonstrated the power of this combination (18): mesenchymal stem cells were transfected with a luciferase vector and labeled with either superparamagnetic iron oxide (SPIO) or Gd for MRI. The authors found that SPIO signal persisted even after complete loss of BLI signal. SPIO signal persisted longer than Gd signal. This demonstrates that simply the presence or absence of signal owing to a contrast agent may not indicate live cells. Often, dead cells and their material (including some imaging agents) can be phagocytosed by macrophages and thus indirectly label these cells. These factors must be taken into account when interpreting in vivo image data.

2.2. Complementary Information

The use of another scientific technique, with a more established label and protocol, is essential to develop and apply a new label and/or imaging modality. For example, histology or microscopy

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Figure 1. Multimodal imaging of labeled human islets. The islets were labeled with a fluorinated emulsion, where the fluorocarbon used also contains a Br atom which allows for Computed Tomography (CT) contrast. (a) Shows $^1$H MRI detection of iron-labeled islets, at 50 islets/ml gel. The dark spots represent single islets (inset). (b) CT image of 10 fluorine-labeled islets. (c) $^{19}$F MR imaging of labeled fluorine-labeled islets, with a phantom containing 10, 50, 100 and 200 islets at high field strength. (d) Lower field strength imaging of fluorine-labeled islets with 500, 1000, 2000, 3000 and 4000 μg of fluorocarbon (top), and 500, 1000, 1500, 2000 and 2500 fluorine-labeled islets (bottom). (e) Ultrasound images of fluorine-labeled islets transplanted intramuscularly in a nude mouse, at 1, 5, 10 and 25 islets per injection. Figure reproduced from Barnett et al. (7), with permission.
has often been used as the ‘established’ technique to confirm in vivo image data. Histology can only be performed at the end-point of an in vivo study, and is thus often supplemented or even replaced by FLI. Thus, most preclinical imaging agents contain a fluorescent component. Confocal microscopy is also typically used to study intracellular localization, although this can also be done with electron microscopy with agents such as iron oxides or quantum dots.

Other combinations of imaging modalities are used to yield different types of information. For example, MRI can be used for localization, together with scintigraphy or SPECT for quantification of cells labeled with both SPIO and radioactive indium isotopes. This has been done clinically to track dendritic cells in melanoma patients (19). More recently, such trials have been combined with PET, as PET can give longitudinal data on cell functionality (20).

The ability of multimodal imaging to yield complementary information is exemplified in a protocol designed to study stem cell implantation (11); BLI was used to assess cell survival, MRI for localization and post-mortem histology to validate the in vivo imaging results. In this example, the cells and the MRI label both had different fluorescent dyes to allow exact localization of the MRI agent using microscopy.

### 2.3. Labeling Strategies

Multimodal imaging requires that cells are labeled with imaging agents detectable by more than one imaging modality. However, this can mean a single agent, or a combination of chemically distinct agents in a cell or a combination of tracing cells that are individually labeled for a single imaging modality (see graphical abstract). All of these combinations have been used in the literature.

#### Table 1. Representative list of the different combinations of in vivo multimodal imaging for cell tracking used in the literature. Agent refers to the type of label used, for example, liposomes or molecular constructs

<table>
<thead>
<tr>
<th>Agent</th>
<th>Cell type</th>
<th>Imaging modalities</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reporter gene</td>
<td>Transfected tumor cell</td>
<td>FLI, MRI, PET</td>
<td>(6)</td>
</tr>
<tr>
<td>Emulsion</td>
<td>Pancreatic islets</td>
<td>FLI, MRI, CT, ultrasound</td>
<td>(7)</td>
</tr>
<tr>
<td>Gold nanoparticles, Gd chelates</td>
<td>Pancreatic islets</td>
<td>MRI, CT, US</td>
<td>(8)</td>
</tr>
<tr>
<td>Iron oxide, PET tracer</td>
<td>Stem cells</td>
<td>MRI, PET</td>
<td>(9)</td>
</tr>
<tr>
<td>MicroRNA-targeted magnetic fluorescence nanoparticles</td>
<td>Neuronal stem cells</td>
<td>MRI, FLI</td>
<td>(10)</td>
</tr>
<tr>
<td>eGFP and luciferase expressing cells labeled with iron oxide</td>
<td>Transgenic murine stromal cells</td>
<td>FLI, BLI, MRI</td>
<td>(11)</td>
</tr>
<tr>
<td>SPIO conjugated to $^{111}$In</td>
<td>Tumor cell line</td>
<td>MRI, SPECT</td>
<td>(12)</td>
</tr>
<tr>
<td>Functionalized silica nanoparticles</td>
<td>Immune cells</td>
<td>FLI, MRI, PET</td>
<td>(13)</td>
</tr>
<tr>
<td>PEGylated complex</td>
<td>Stem cells</td>
<td>MRI, luminescence</td>
<td>(14)</td>
</tr>
<tr>
<td>Molecular complex</td>
<td></td>
<td>MRI, CT, luminescence</td>
<td>(15)</td>
</tr>
<tr>
<td>Particles</td>
<td></td>
<td>BLI, PET, MRI, FLI</td>
<td>(16)</td>
</tr>
<tr>
<td>Antibodies</td>
<td></td>
<td>BLI, scintigraphy</td>
<td>(17)</td>
</tr>
</tbody>
</table>

MRI, magnetic resonance imaging; PET, positron emission tomography; FLI, fluorescence imaging; BLI, bioluminescence; SPECT, single photo emission computed tomography.

#### Table 2. A general summary of the main issues that arise with multimodal imaging for cell tracking, and their common solutions

<table>
<thead>
<tr>
<th>Issue</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imaging often detects only the presence of label, whether or not it is in the relevant cells</td>
<td>Reporter genes, or coupling to a second imaging agent that is known to be cell-specific</td>
</tr>
<tr>
<td>Effect of label on cells, particularly long-lived cells</td>
<td>More extensive testing of labeled cells, including specific functionality assays</td>
</tr>
<tr>
<td>Difficulties in image registration with different imaging modalities, particularly with deformable tissues</td>
<td>Various strategies have been developed, such as subject immobilization, addition of references, computer algorithms, and the use of hybrid scanners</td>
</tr>
<tr>
<td>Cost and increased imaging time</td>
<td>The introduction of hybrid scanners can reduce imaging time and cost</td>
</tr>
<tr>
<td>Discrepancy between data from different imaging modalities</td>
<td>These typically arise owing to differences in sensitivity to cell viability or functionality between the labels</td>
</tr>
<tr>
<td>Multiple and longer imaging sessions, and increased and more frequent use of anesthetics in preclinical studies</td>
<td>Hybrid scanners where possible, and careful planning is necessary</td>
</tr>
<tr>
<td>Complex anatomy and organ movement (bone, bone marrow, lymph nodes, heart, lung)</td>
<td>Specific adjustments and data processing for recording and overlaying data.</td>
</tr>
</tbody>
</table>
When a combination of cells with individual labels is used, it is assumed that the cells will colocalize and that the different labels do not affect the cells differently. Furthermore, it is possible that label is not taken up uniformly within a cell population, generating mixed results in viability or functionality assays (21), and the possibility that the image data does not reflect the behavior of the majority of nonlabeled cells. Lastly, inadvertent label transfer may occur to nonrelevant cells.

A final factor that can affect image interpretation is the dilution of label with cell division. This will not occur if the cells are labeled in situ, or if a reporter gene is used. However, even with rapidly dividing cells, the number of cell divisions that may occur result in tolerable quantification errors (22). All of these issues must be considered when designing the experimental model.

2.4. Effects on Cells

A recent paper (23) discussed the confounding effect that the simple addition of biomaterials, such as contrast agent nanoparticles, can have on standard cytotoxicity assays. This effect occurs owing to the formation of a coating of biomolecules, particularly proteins, on the nanoparticle surface (24), significantly altering the cell medium. The simple aggregation and sinking of label particles can also affect the actual concentration to which the cells are exposed. Other key factors that must be considered are the lifetime of the cell vs the lifetime of the label, viability, in vivo toxicity and clearance and specific effects on cell functionality. Most often the concentration of label added to cells is selected to balance any effects on the cell with maximal label loading.

In addition to the general effects of adding agents, particularly nanoparticles, to cells, specific functional effects can also occur. Several studies have looked into the effect of labeling various cell types with iron oxide agents for MRI, with mixed results on cell viability and functionality. For example, labeling human embryonic stem cells with SPIO had no effect on pluripotency or differentiation capacity compared with nonlabeled controls (25). In this study the amount of iron loading was about 4.5 pg/cell, which is a typical value although the authors used a shorter incubation time and lower SPIO concentration when labeling. The label persisted for 21 days in the cells. Other studies found that labeling with SPIO affected the insulin synthesis in a pancreatic cell line (26), cytokine secretion in macrophages (27) and mobility of neural stem cells (28,29).

However, the majority of published data show minimal or no significant effects of labeling with iron oxide on cells, at least on standard assays.

With other imaging modalities, particularly SPECT and PET, the effects of radiation from the agent must also be considered, particularly when labeling long-lived cells such as stem cells (30,31). This is in addition to limitations in the detection time frame owing to the radioactive half-life of the isotope used.

2.5. Multimodal Imaging in Cellular Therapy

The role of imaging in cellular therapy has been reviewed elsewhere (1). Clinical cell tracking has generally been done using scintigraphy and SPECT, and more recently with MRI using iron-based labels, although this may be hindered by the recent removal of several MRI contrast agents from the market. FLI, which is the most commonly used ‘second’ imaging modality in the preclinical world, has limited applicability in humans owing to penetration depth issues. Furthermore, increased cost, imaging time, toxicity and side-effects, the need for more trained personnel and simply the novelty of the techniques has somewhat limited the use of multimodal imaging in clinical cellular therapy.

However, the multitude of preclinical studies now available has demonstrated the necessity for multimodal imaging in optimizing cellular therapy, especially as the field of cellular therapy itself develops and becomes more complex. Figure 1 shows an example of a multimodal imaging of labeled pancreatic islets.

3. CONSIDERATIONS

Several assumptions are typically made when imaging labeled cells for in vivo imaging. These are summarized in Table 1, along with the most common strategies used to counter them.

Some agents have been developed and shown to be chemically feasible but have not been tested in vivo or even in cells (see Table 2 for examples). Such testing is especially important when the agents contain heavy metals, which are extremely toxic if released from stabilizing chelates. For example, Gd chelates, commonly used as blood pool agents for MRI contrast, are now indicated as a risk factor for systemic nephrogenic fibrosis (32). Carbon nanotubes and silica nanoparticles, both of which show great promise as preclinical imaging agents, may also have issues owing to long retention times (33).

4. SUMMARY AND OUTLOOK

The use of reporter genes revolutionized fluorescence microscopy, as it allows the production of reporters such as a fluorescent protein linked to another gene product or constitutively expressed. Thus, the presence of the reporter product is highly restricted to a specific cell type, is not diluted by cell division (if the gene becomes integrated) and occurs only in viable cells. Luciferase is typically used as a reporter gene product for BLI, and reporter genes have also been developed for MRI. These reporters are typically proteins involved in cellular metal homeostasis, particularly iron storage (34). A recent study used the expression of a metabolically biotinylated luciferase that is membrane bound (6). This construct allowed in vivo BLI, and also targeting of other imaging agents to the membrane-bound biotin using streptavidin-bound Alexa750 (FLI), 111In (SPECT) and magnetic nanoparticles (MRI). This approach couples targeting and noninvasive imaging. Such an approach is easily amenable to multimodal imaging, but is restricted to genetically modified cells and subjects on biotin-deficient diets. The use of reporter genes for imaging may overcome some of the disadvantages of conventional labels, at least in preclinical models.

It is vital to consider what information is required from the images (localization, viability, functionality or quantification) in order to determine which imaging modality would be most applicable, and if multimodal imaging provides relevant, non-redundant information. The practicality of an imaging scheme must also be considered, in terms of the number of imaging sessions required, cost, imaging time and (repeated, frequent) anesthetic use for preclinical studies. A multimodal imaging study can yield more information per subject, but requires more and longer imaging sessions per subject. For example, simply organizing the logistics can become difficult: in one paper (20), patients received injections of a radioactive tracer for PET after receiving cellular therapy for cancer. The PET tracer
accumulated in treated lymph nodes, but this presented new problems as some of these patients were also scheduled for lymph node resections, thus potentially exposing the surgical staff to the residual radioactivity. Furthermore, the PET tracer itself had to be prepared and transported from the cyclotron in time for the cell injections, within the usable lifetime of the tracer. These kinds of studies require precise and thorough planning, and can involve over 50 people in the clinic. Despite these issues, PET is a common clinical imaging modality that is now more often being combined with MRI (and has long been combined with CT).

Finally, the addition of specific functionalities to multimodal imaging agents is under development. For example, one agent consists of a iron oxide core with quantum dots functionalized with an anti-cancer agent and a targeting motif (35). In this agent, the quantum dots remain quenched until activated by specific intracellular reactions triggered by drug uptake. Overall, despite the feasibility challenges, in our opinion, multimodal imaging will offer many advantages to guide progress in cellular therapeutics.

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