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MISSION STATEMENT

The International Society for Strategic Studies in Radiology is a non-political, not-for-profit organization whose purpose is to define and investigate strategic, scientific and economic issues of global importance to the field of radiology. The Society aims to foresee national and global developments in radiology and allied fields, and to strategically communicate important issues to radiologists, physicians in other medical fields, industry and governments. By forming a strategic partnership with industry in order to define important and relevant fields of cooperation and common engagement for the best possible cooperation and successful future of both partners, IS³R hopes to envision schema for actively influencing research, clinical practice and teaching now and in the future, and to prepare radiologists, their partners, and their clients for future developments. The Society aims to influence the flow of capital and investment into the most scientific and strategic fields, and to influence health care politics for the benefit of radiology, its patients and customers.
# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTRODUCTION</td>
<td>5</td>
</tr>
<tr>
<td>INTERVENTIONAL RADIOLOGY: NEW FIELDS AND CHALLENGES</td>
<td>7</td>
</tr>
<tr>
<td>QUALITY &amp; SAFETY</td>
<td>10</td>
</tr>
<tr>
<td>MOLECULAR IMAGING &amp; NANOTECHNOLOGY</td>
<td>13</td>
</tr>
<tr>
<td>REGULATORY HURDLES</td>
<td>15</td>
</tr>
<tr>
<td>REFERENCES</td>
<td>18</td>
</tr>
</tbody>
</table>
INTRODUCTION

The modern transformation of radiology into an essential guiding hand of medical practice began with the invention of computed tomography imaging in the early 1970s and continues today with novel applications based on molecular methods and nanotechnology. The topical agenda of the 7th Biannual Symposium of the IS³R, “Advancing Radiology Through Informed Leadership,” was chosen by Program Committee Chairman Maximilian Reiser to promote exploration and discussion of radiology’s continuing transformation and to further explore its implications for the care of patients and, more broadly, its impact on the health care systems of the world.

Major themes of the Symposium included the increasing role of image-guided therapeutic interventions, the central role of quality and safety, the emergence of molecular imaging and nanotechnology and new paradigms for patient management related to cardiac, oncologic and emergency imaging. Regulatory hurdles for technical innovation were addressed for both drugs and devices.

The IS³R brings together thought leaders from academia and industry from around the world to share ideas, points of view and new knowledge. The following summary reflects the contributions of these diverse voices and provides a unique window for looking at important trends that are shaping the future of radiology.
The technology for minimally invasive, image-guided interventions is advancing rapidly, aided by the convergence of innovations from numerous scientific and medical disciplines. As a result, a host of new options for the minimally invasive diagnosis and treatment of disease are broadening the horizons of interventional radiology (IR).

Fusion of images from different radiological techniques is allowing more accurate and targeted intraoperative navigation. For example, pre-operative 3D CT angiography is being successfully co-registered with intraoperative 2D angiography to provide better depth perception and navigation during angiographic liver interventions, with motion correction for breathing (1). Similarly, preoperative 3D CT is being fused with intraoperative ultrasound. Anatomical imaging is also being combined with functional or molecular imaging. Thus the use of ultrasound to identify suspicious nodules may be immediately followed by the use of a gamma probe to determine whether the nodules display functional activity indicative of cancer (2). In patients suspected to have prostate cancer, targeted biopsies guided by the fusion of ultrasound with MRI and MR spectroscopy are allowing better cancer detection (3).

“Smart” devices are also improving navigation during image-guided interventions. For example, the insertion of miniature “smart” electromagnetic sensors into interventional tools such as guide wires or needles allows their progress through the body to be followed on a GPS-like tracking system (4). In this approach, the magnetic space of the patient is co-registered with preoperative 3D imaging data, essentially creating a road-map for the intervention. Not only does this lessen the need for intraoperative imaging (which may involve exposure to ionizing radiation and/or contrast media), it also opens up more possibilities for bringing information from sophisticated preoperative imaging techniques—such as PET or MRI—directly into the interventional process (4). Robotic instruments for needle placement are now available that are relatively inexpensive. CT-integrated robots place needles more accurately and consistently than humans, and with the use of treatment planning software, they can identify and access the most direct routes to their targets. Although their cost-effectiveness has not yet been proven in large-scale studies, they work rapidly and precisely, and they also reduce interventional radiologists’ exposure to radiation (5).

Further tailoring of interventions is also becoming possible through the advent of “smart” drugs that are specifically designed to complement interventional techniques (6). One example is Thermodox™, which consists of liposomal capsules that contain the chemotherapeutic agent doxorubicin. The capsules release the doxorubicin only under the high temperatures induced by thermal ablation. When the capsules are injected into the tumor vasculature at the time of ablation, the doxorubicin takes effect throughout the tumor, including at the edges, where thermal ablation alone tends to be ineffective. Because the doxorubicin is released only in the heated area, its toxic side effects are limited. Another concept consists in combining the application of liposomal capsules with regional hyperthermia, eg, with MR-guided focused ultrasound (7).

With advances in nanotechnology, the potential for the development of “smart” drugs appears limitless. For instance, targeting antibodies or ligands could be attached to drug-carrying capsules made with bound metal (to allow imaging). Not only the ligands, but the agents inside the capsules could be varied to target specific disease characteristics. In a new paradigm of “molecular IR,” molecular imaging-guided interventions could be used to determine the molecular characteristics of a disease and to subsequently deliver the appropriate targeted therapy (7).
Emerging cellular therapies could also play a role in IR. Preliminary research suggests that the insertion of autologous bone marrow cells after myocardial infarction can improve left ventricular function and myocardial viability (8). Various imaging techniques, including MRI, PET and SPECT, have been used to assess the functional effects of cell therapy, and methods for tracking and labeling cells with radionuclides and superparamagnetic agents are being developed (9). Although extensive clinical trials are still needed, cellular therapy could potentially become an important—and more widely available—alternative to heart transplantation for the vast numbers of patients suffering from severe heart disease (10).

MR-guided focused ultrasound (MRgFUS) is another novel interventional technique that could have applications in a multitude of patients. In this procedure, a high-intensity ultrasound beam (5000-10,000 times stronger than the ultrasound beams used in diagnostic radiology) is directed at a focal point, where it causes intensive local heating and tissue destruction. No incisions of any kind are made in the patient, and the entire procedure, which is performed on an outpatient basis, takes place inside the bore of the MRI magnet. MR imaging is used to guide the beam, avoiding critical structures such as nerves, scar tissue, surgical clips, and bowels, and MR thermal mapping is used to monitor the temperature rise (through a phase shift) and tissue destruction (11). To date, more than 3,000 patients have received MRgFUS for uterine fibroids (12). In a study conducted in the UK, 80/100 patients treated with MRgFUS for uterine fibroids had significantly less severe symptoms 6 months after treatment (13). Data suggests that MRgFUS may have more long-lasting effects in the treatment of uterine fibroids than the more invasive alternative of myomectomy. MRgFUS has also been used for the palliative treatment of bone metastases. Destruction of periosteal nerves has been suggested as a possible explanation for the analgesic effect, which cannot be explained simply by the avid absorption of ultrasound waves by bone.

Although the inability of ultrasound waves to pass beyond bones limits the areas that can be directly accessed by the technique, MRgFUS is now being evaluated for the treatment of primary liver tumors. Although 90% of the energy is reflected back by the calvarium, enough power remains to thermally ablate a glioblastoma multiforme or liquefy a clot in acute stroke. (The reflected energy is absorbed by a waterbag). Moreover, results in the treatment of breast cancer as well as soft-tissue metastases using MRgFUS have been reported (11). As a non-invasive, imaging-controlled treatment, MRgFUS has immense potential for the safe and cost-effective treatment of a wide variety of conditions. Radiologists have an essential role to play in the development and application of this technique, as it requires substantial skill in the use of imaging to guide the ultrasound beam.

The developments described above suggest great potential for the growth of IR. However, despite a history of introducing innovative and widely applicable techniques, the field of IR has grown at a very slow rate, such that in 2002 there were only around 200 IR training positions in the United States (as compared to around 2300 in cardiology the year before) (15). To some leaders in the field, these numbers suggest that the very survival of IR is at risk (16).

The limited growth of IR is related to its status as a subspecialty of radiology, in which the focus is mainly on diagnosis, with little or no patient contact. As a rule, patients are referred to IR from organ-based specialties—e.g., urology or cardiology—which remain responsible for the patients’ overall care. Repeatedly, techniques that have started out in IR have ended up being adopted by organ-based specialties, and the referrals to IR for these procedures have dropped off. For example, by 2003, cardiologists in the US had caught up with radiologists in the performance of peripheral vascular stent procedures, and now the majority of such procedures are performed by cardiologists (17). With their direct access to patients, their comparatively large numbers of personnel,
and their greater financial resources and research activity, organ-based specialties have tremendous advantages over IR in the competition for patients (18).

What can be done to sustain and enhance the relevance of IR? One advantage that IR may have over competing fields is in the area of skills—particularly in the use of advanced imaging equipment. To build on this advantage, adjustments to the standard IR training may be needed. Arguably, more of the training could be focused on IR procedures and equipment and less on diagnostic imaging, while still providing the core skill necessary to achieve Board certification in radiology. Clinical skills should be emphasized. Efforts should also be made to increase recruitment into IR training programs, perhaps by looking for candidates outside the field of radiology, in specialties that attract more physicians who seek patient contact (19). The move in Europe to a 3-year core training with 2 years devoted to sub-specialty training should help in this regard (20).

The relationship of IR to radiology may need to be reconsidered. Should IR be separated from radiology and become an independent clinical specialty? Should it remain within radiology? Or should it be organized into partnerships with separate organ-based specialties? Each scenario has advantages and disadvantages (21). One advantage of remaining within radiology is the access this affords to advanced imaging equipment—access that is not as readily available to physicians in organ-based specialties. However, if IR is to stay within radiology, the differences between the work of interventional radiologists and that of purely diagnostic radiologists must be acknowledged and addressed. For example, since the former cannot bill as many procedures as the latter, they should perhaps have a separate accounting system. More hospital and departmental resources should be devoted to marketing and community outreach for IR. Furthermore, regardless of which organizational approach is pursued, interventional radiologists should seek to obtain the status of true clinicians, with admitting privileges and the administrative support and infrastructure that will allow them to take more direct responsibility for patient care. Interdisciplinary collaboration and involvement in research are also essential to enhance the growth of IR. At present, IR lags far behind organ-based specialties in the publication of original research. Many promising new interventional techniques (including most of those described above) have yet to be validated through extensive clinical trials. More studies are needed to demonstrate their benefits to patient outcomes (13) as well as to workflow. Furthermore, with the continual advancement of technology, there are endless possibilities for developing new interventional approaches. To enhance their contributions to both routine care and research, interventional radiologists should build strong collaborative ties with physicians in other specialties as well as with the radiochemists, physicists, computer scientists, engineers and basic scientists whose knowledge is essential for the development of new minimally invasive techniques. One thing is certain: Regardless of whether they continue to be applied under the rubric of IR, the skills of interventional radiologists are going to be increasingly in demand (23). Furthermore, the interventional radiologist who can deliver a comprehensive range of emergency procedures will continue to be held in high regard by professional colleagues.
QUALITY & SAFETY

Rising healthcare costs in the developed world have led to increased scrutiny of the quality of care being purchased. Four broad components of quality healthcare are now widely recognized: appropriateness of care (as opposed to overuse, underuse or misuse of procedures); coordinated care that links providers and information; safe, error-free care; and patient-centered, timely care (24). The methods for evaluating these aspects of quality, however, are still being developed and debated.

One common way of gauging quality has been to compare usage of medical procedures across geographic regions, institutions and individual providers. For years, wide variations in usage have been acknowledged to exist in the United States (23). However, solutions to this problem have been slow in coming. As recently as 2003, McGlynn et al. found that on average, Americans received only about half of recommended medical care (as defined by a panel of physicians working on the study) (26), with the percentage of recommended care received varying substantially by medical condition. Interestingly, the latter study found that underuse of medical processes was a much greater problem than overuse, with 46.3% of study participants not receiving recommended care and 11.3% receiving superfluous and potentially harmful care. In radiology, it appears that the opposite may be true. A study of the healthcare provided to employees of General Electric suggested that approximately half ($60 million-worth) of the radiological tests ordered were unnecessary (27). Imaging costs are accounting for a growing share of hospital expenditures (27). Nevertheless, when used appropriately radiological services can make healthcare better, quicker and cheaper, most often by diminishing diagnostic uncertainty and obviating the need for additional (and more invasive) procedures. More studies are needed to demonstrate that radiology can improve outcomes and reduce costs at every level, from the clinician, to the department, the hospital, the insurer, the community, and the nation (28,29).

Refinements in the criteria for test usage and reporting can lead to considerable increases in cost-effectiveness. For example, a study in the UK found that the incremental cost of using breast MRI would be approximately $56,153 per cancer detected (30) if it were applied in all patients who underwent mammography, but only about $23,286 if it were applied in place of mammography for patients with BRCA1 and approximately $30,374 if it were applied in addition to mammography in patients with BRCA2 (30). Partly as a result of such findings, the use of MRI in patients at high risk for breast cancer is now being funded by the National Health Service (NHS) (31). Another study found that by not reporting lesions smaller than 6 mm seen on CT colonography, radiologists could substantially reduce costs and complications while producing only a trivial decrease in cancer detection (32). To help develop and improve imaging recommendations, more studies are needed that focus on whether imaging tests are likely to improve clinical decision-making and hence clinical outcomes. Modeling can be used to estimate the impact of radiological tests on clinical outcomes. To yield results that are applicable to a broad spectrum of clinical settings, more meta-analyses of small studies are needed, as well as more large, multicenter studies—though it is important not to conduct the latter too early, before the technology and radiologists’ skills have matured. Aids for designing and implementing clinical trials of imaging are readily available. For example, the Web site of the American College of Radiology Imaging Network provides access to standard enrollment form templates and sample protocols (33).

Decisions about cost-effective radiological test usage may be complicated by concerns about exposure to ionizing radiation. Increasingly, patients are becoming aware of the risks of radiation exposure, and organized efforts are being made to address the matter. In the late 1990s, the European Community
issued Directive 97/43/EURATOM outlining the need for justification, optimization, and auditing of the use of ionizing radiation in healthcare. Over the past 5 years, the European Commission issued European Guidelines for Multislice Computed Tomography (an update of an earlier set of guidelines for the use of CT) (34) and launched the project “Safety and Efficacy of Computed Tomography: A Broad Perspective.” The latter project brings together radiologists and physicists from leading European academic institutions to study CT justification and radiation dose optimization and measurement and to design practical tools for improving practice (e.g. clinical decision trees and optimized CT techniques). The project has resulted in published papers, conference presentations, and a report to the Commission and has had an impact on patient care. However, dissemination of the information has been incomplete. Research is being outpaced by advances in imaging technology. Furthermore, as CT is now widely available, relatively affordable and quick, guidelines for limiting its use are being ignored to meet the demand for imaging, and it appears that many, perhaps even most, CT examinations performed in Europe are being requested in advance of patients’ needs (35). Increasing the availability of both ultrasound and MRI will reduce reliance upon techniques involving x-rays, particularly for young patients at higher risk (36). To address the problem of radiation exposure more effectively, the government must continue to support research and not only pass but enforce laws through auditing; researchers must work closely with industry to ensure that quality and safety measures are incorporated into technology development; and leaders in government, healthcare and industry must change attitudes by prioritizing quality and long-term safety above (real or imagined) productivity.

Various approaches are being pursued for improving—and not just retrospectively assessing—the quality and appropriateness of healthcare. For patients on Medicare in the US, efficiency measures have been proposed for the use of certain radiological examinations, including MRI of the lumbar spine and mammography. Recall rates after mammograms are now being evaluated to determine whether mammograms are being performed in the appropriate patients (37). Meanwhile, private insurers have begun to scrutinize utilization rates per thousand patients. Such measures may only just begin to address the problem. Ultimately, to bring about changes in practice, information on performance at all levels will need to be made routinely available—a process that will require substantial changes in health information systems, including automated entry of essential data for clinical decision-making (24). This will help to establish baselines and standards of care. Accreditation systems offer an important means of assessing and raising the quality of healthcare services. Healthcare accreditation systems are now in use in the US, Finland and Australia. In 2006, the Royal College of Radiologists (RCR) and the Society and College of Radiographers began to develop the Radiology Accreditation Project (RAP) to assess radiological services in the UK (38). Accreditation by the program requires demonstration of appropriate protocols for monitoring and assuring the quality of service relative to published standards for safety, patient experience, clinical performance, and utilization of resources and workforce. In addition, it requires the development of a quality improvement plan with realistic, achievable goals. The process involves assessment by independent auditors as well as self-assessment. At present, the RAP is still in the pilot stage and is voluntary. However, it is hoped that by providing information about the strengths and weaknesses of radiological services and the improvements to which they are committed, it will eventually offer a basis for making comparisons between service providers.

The RCR is also working to establish a system for relicensing and re-certification of radiologists in the UK, with the goal of identifying poorly performing ones. It is known that the performance of approximately 2% of radiologists in the UK falls 2 standard deviations below the mean, yet only 0.5-1% of radiologists are referred for evaluation by the National Clinical Assessment Service (39). Measuring the overall
performance of individual radiologists can be a tricky task. Theoretical knowledge can easily be assessed by examinations, while practical skills can be assessed, though perhaps somewhat less reliably, with more complex tests. However, assessment of professionalism and behavior has generally relied on informal feedback, which is often considered unreliable and ignored. In a survey that asked RCR members and fellows to choose the best methods for identifying poorly performing radiologists, third place went to peer review of cases, and second place went jointly to attendance at discrepancy/error/complications meetings and on-line assessments linked with educational programs. First place went to a less conventional solution: multiple-source feedback, consisting of a confidential questionnaire for peers and patients, ranking of behavior, skills and knowledge, and the assignment of scores for comparison with those of other radiologists and doctors, ideally both nationally and locally. The proposed multi-source feedback approach does not involve the assignment of pass or fail marks, but instead requires further evaluation of any radiologists who receive consistently low scores. Perhaps this approach won out because it was perceived as allowing a fairer and more nuanced evaluation, more reflective of the radiologist’s overall performance and of the context in which he or she works. Although much emphasis is now being placed on increasing efficiency, it is important to keep the focus of attention on the patient. Improving the quality and safety of care will in turn improve productivity. In other words, business objectives and patient care objectives do not conflict. For example, the business objective of an inpatient unit is to reduce the length of the average patient stay in the hospital; for radiology, this means optimizing the turnaround time from study request to study completion, which both shortens the patient’s stay and improves the patient’s care (40). The use of flow charts in work process analyses can help identify ways to reduce waste and the potential for error. Some of the solutions may be non-intuitive. For example, net savings may result from hiring an extra full-time technician to keep an expensive piece of equipment (e.g., an MRI machine) in operation during examination preparation times and the primary technician’s lunch hour and coffee breaks. Conversely, for equipment that is less expensive to purchase and operate, it may be more cost-effective to buy many units, so that they can readily be used whenever needed, even if they are not constantly in use. Improving productivity and care requires establishing the right balance between equipment and staff and providing the necessary training for staff to make optimal use of equipment (41). At present, the error rate in the U.S. healthcare industry far exceeds those in the airline, telecommunications, and computer industries. Teamwork and communication are essential for improving this state of affairs. Traditionally, the culture in medicine has encouraged physician autonomy over teamwork. However, physicians and staff with average skills but an exceptionally well-designed work process will outperform those with better skills but a less coordinated work process (42). Approaches used to encourage communication in the airline industry, such as anonymous reporting of errors and elimination of penalties for errors, have been applied in healthcare organizations and have been found to increase the rate of error reporting, stimulate discussions to improve work processes and ultimately decrease the incidence of errors (43). Multidisciplinary team meetings allow radiologists to make recommendations for the clinically relevant use of imaging and deliver clinically relevant interpretations. Communication amongst physicians and staff during the delivery of care is essential to prevent errors. For example, it is the responsibility of the radiologist to speak up if a radiological test appears to have been requested unnecessarily or if the use of contraindicated contrast media has been ordered.

Information technology, such as decision support for order entry and computer-aided diagnosis algorithms, can also help reduce errors and improve the quality of care. Furthermore, teleradiology is increasingly being used around the world to augment local radiological services (44,45). In the US, the most common use
of teleradiology has been to allow radiologists to interpret images from home, but approximately 50% of radiological practices are now using outside service providers for interpretations on nights and weekends. In parts of Europe and Asia, teleradiology is being used across borders either to serve areas with a shortage of radiologists or to allow secondary consultations with specialists, or both (46). The use of Indian teleradiology services in Singapore has improved care by reducing patient wait times and allowing more rapid clinical decision-making; it has also forced radiologists in Singapore to work faster in order to prove their worth.

Limited communication may be the most significant drawback of teleradiology, as direct access to complete clinical information, including prior imaging studies, is rarely available, and teleradiology reports are usually submitted by e-mail or fax, without direct contact between the radiologist and the referring physician, who may be unfamiliar with the radiologist’s reporting style. In addition, auditing is needed to ensure that teleradiologists meet the same certification and training requirements as local radiologists and that adequate clinical and technical standards are being maintained. Although a study comparing interpretations made by NHS radiologists with those made by teleradiologists at an independent service provider found few discrepancies, most of the cases examined were of low complexity (47). The RCR, the American College of Radiology and the European Society of Radiology (in conjunction with the European Union of Medical Specialists) have all issued fairly similar guidelines for quality assurance in teleradiology (48).

MOLECULAR IMAGING & NANOTECHNOLOGY

The potential of molecular imaging to revolutionize healthcare is often spoken of in rhapsodic terms. Yet the practice of molecular imaging (MI) within radiology continues to be confined to a relatively small number of institutions and settings, and the reality is beginning to catch up with the hype surrounding the field. Though MI already has clear applications in neurodegenerative conditions, cardiovascular disease, inflammation (49), and above all, cancer, a number of obstacles are impeding progress toward its broad clinical implementation—some of them avoidable, others less so.

It is hoped that “system characterization” by imaging probes will ultimately allow molecular knowledge developed in the laboratory to be translated to clinical disease management (50). In combination with in-vitro molecular markers, MI (performed mainly with multimodality techniques such as PET/CT and PET/MR) (51) should facilitate earlier, even preclinical disease detection and characterization, prediction of prognosis, targeted treatment, and precise treatment follow up, allowing “personalized medicine” that is tailored closely to the individual patient. However, both physical and technical factors make achieving this vision a highly complex and challenging endeavor. The body is equipped with many barriers to molecular transport—such as nuclear, cellular and vessel walls, and the blood brain barrier. The development of probes suitable for use in humans is limited by pharmacokinetics, biocompatibility and toxicity concerns, and imaging speed and resolution. Contrast agents and radiopharmaceuticals must have sufficient half lives and must aggregate in sufficient concentrations to allow effective imaging once they have reached their target destinations. In addition, for
any one disease, a limited number of overexpressed molecular markers will exist for targeting (52).

Molecular interactions and disease processes are complex, and knowledge of them remains limited. Although molecular imaging with PET has been used successfully to explore how new drugs interact with targeted systems and to narrow the feasible dose ranges, PET imaging results can also be misleading. For example, during its development the anti-psychotic drug aripiprazole had no clinical effectiveness at the feasible dose range identified by PET, but in practice, the drug was found to be effective and safe at substantially higher doses. Perhaps the tracer used to image the drug did not have a sufficiently long half-life, or perhaps the mechanism of the drug was not as pure as anticipated. While molecular imaging can show on-target biological activity and provide essential information about the biodistribution of drugs, both more knowledge and more probes are needed to increase the reliability of molecular imaging for assessing drug effects. In certain instances, it may soon be possible for MI to play a role in disease stratification, patient selection for therapeutic agents, and dose selection (52), but MI is unlikely to replace clinical endpoints in drug development for a long time to come.

Nanotechnology offers enormous versatility in the development of MI probes. The advent of particle replication in non-wetting templates (PRINT) technology has made it possible to mold nanoparticles small enough to cross the blood-brain barrier (53). Not only the size, but the ability of nanoparticles to deform can be tailored to their purpose, increasing or limiting their uptake. In addition, nanoparticles can be filled with cargoes (e.g., antibodies) and can have entities (e.g., targeting ligands) attached to their surfaces (54). Applications for which nanoparticle-based contrast agents are expected to be introduced into clinical practice within the next 10 years include imaging of metastatic lymph nodes, vulnerable atherosclerotic plaque imaging, macrophage imaging for inflammatory and degenerative diseases (e.g., multiple sclerosis, polyarthritis and osteomyelitis), and imaging of amyloid plaques in Alzheimer’s disease. Many more probes might now be on the horizon were it not for the vast amounts of time and money required to overcome regulatory hurdles. In the US, this problem has been exacerbated by misconceptions about regulatory requirements. Many applications for pharmacology/toxicology studies have been submitted with much more background data than necessary (55). In 2006, the FDA issued guidelines for exploratory investigational new drug (IND) studies (e.g., screening studies or microdose studies) that are conducted early in phase 1 of the trial process, involve very limited human exposure for a short time period (e.g., 7 days) and have no diagnostic intent (56). The guidelines explain that because exploratory IND studies present fewer potential risks than do traditional phase 1 studies that look for dose-limiting toxicities, they can be initiated with less, or different, preclinical support than is required for traditional IND studies (56). Exploratory IND studies allow a large number of agents to be tested rapidly in order to distinguish the promising from the not-so-promising, decreasing the likelihood of failure in later (and more costly) clinical trials.

While exploratory IND studies are extremely useful, further simplification of regulatory processes is necessary, as is the creation of a viable business model for probe development. The development of MI probes that deliver drugs could allow much more targeted treatments of disease with less morbidity. However, trials of such probes are especially complex and doubly risky for the sponsors, since either the imaging probe or the drug may fail. Collaboration between academia and industry in probe development has been hampered by concerns about intellectual property rights. Furthermore, lack of reimbursement discourages experimental use of molecular imaging in hospital settings, particularly since the required multimodality imaging techniques are expensive (57). Despite these obstacles, collaboration between industry and academia is possible in MI research and probably essential for academic institutions. In a recent survey of academic radiology departments with MI programs in the US, only 16.1% had no
alliances with industry (58). Another barrier to progress in the clinical implementation of MI is the need for expertise from multiple disciplines. As it is, radiologists tend to have little understanding of molecular biology, and few have any familiarity with molecular imaging, since it is not included in standard clinical training. To encourage more radiologists (and other physicians) to become involved in the field, MI rotations should be created and more MI fellowships should be offered. In the survey referenced above, respondents identified staff training and recruitment as the most essential elements for success (58). In light of the extensive training in both basic science and multimodality imaging required for the field, the creation of a new specialty of MI may be warranted, with the possibility of dual boarding in diagnostic radiology and MI, or nuclear medicine and MI.

Because MI involves not only multiple disciplines but multiple kinds of technology, its success will require major investments in human resources and infrastructure. It will also require significant changes in organizational structures and the setting aside of turf battles. Conceivably, molecular biology, pathology and imaging could be combined under one umbrella, perhaps that of “biomedical imaging” or “imaging science” (60). Alternatively (or concomitantly), the typical “Department of Diagnostic Radiology” could be replaced by a “Department of Diagnostics,” combining pathology with radiology and MI, and using pathological analysis of genes and proteins to identify targets and create agents for MI in individual patients. Regardless of the specific restructuring plan chosen, broad clinical implementation of MI will depend on collaboration, co-training, and co-ownership.

REGULATORY HURDLES

Approval hurdles for pharmaceutical media and technology

Regulatory requirements pose daunting barriers to the introduction of radiological innovations. For example, a pharmaceutical company is likely to spend over $300 million to bring a new contrast agent to market, in a process that typically takes about 9 years. (The cost for bringing a therapeutic agent to market is now approaching $1B!). And that is only the beginning. For each new indication, about $15 million more in funds is required and around 5 more years although most radiologists ignore the package insert and engage in “off-label” use. While the process is somewhat less onerous in Europe than in the US, regulatory hurdles appear to be increasing around the world.

The need to meet different regulatory requirements in different areas of the world adds to the complexity and costs of the process. Currently, regulations differ in the EU and the US, China and Japan, with the regulatory policies of the latter two countries being closer to those of the US. In both the EU and the US, regulators look for accuracy (e.g., as measured by sensitivity and specificity), reliability, clinical value and a favorable risk/benefit ratio. However, far more contrast agents have been approved in Europe than in the US to date. This may be partly because US regulators seem to place a greater emphasis on the demonstration of p-values showing statistically significant increases in value over existing technology or agents, whereas EU regulators are more content to see any demonstration of utility, even if it is only for a secondary aim (61). Furthermore, in the US, the requirements are often not aligned with the intended use of the agent or device. For example, even if the manufacturer’s goal for a contrast agent is to increase tumor visualization, the FDA may demand that the manufacturer also demonstrate an improvement in
tumor diagnosis. Such a demand may prevent a useful agent from coming to market, or may require it to be tested (and approved) for a narrower cohort and indication.

Irrational elements in the approval process only add to unavoidable difficulties already in place. Advances in technology occur faster than does the development of new contrast agents. Thus delays in the approval process for an agent can make the agent or the application tested obsolete by the time it becomes available. Moreover, partly because of the fast pace of technological change, standardization of imaging protocols in multi-center trials is difficult, as is comparison of the results of multiple single-center studies. For both devices and pharmaceuticals, even clinical trial protocols may be outdated before the trials are begun, let alone completed.

Interestingly, the faster technology advances, the slower the approval process seems to become. For example, in 2005 it took 2 months for one manufacturer to obtain approval of computer-aided diagnosis (CAD) for virtual colonoscopy in the US, but the same manufacturer has since spent more than 24 months working toward approval of CAD for other applications, such as in the breast and lung. In a phenomenon sometimes referred to as regulatory “creep,” the FDA has repeatedly come back to manufacturers and pharmaceutical companies with requests for more data, more details, and stricter adherence to protocols. To prevent this, the FDA should be required to make binding agreements about what is required at the beginning of the regulatory process. Recently, the FDA began to call for clinical trials to assess the use of a contrast agent on a new imaging platform, even when the contrast agent has already been approved. This shortens the effective life-time of the new device, reducing the incentive for the manufacturer to invest in the trial process.

In Japan, regulatory requirements have also multiplied recently. According to a report prepared by the United States International Trade Commission (Washington, DC), the average approval time for a new medical device in Japan in 2004 was 1083 days, compared to 356 in the United States (63). Apparently as a result, the number of new device applications submitted by US companies for review by Japanese regulators fell dramatically, from 132 in 2003 to only 8 in 2005—a decrease of 94%. Representatives of the US medical technology industry cited “burdensome applications and an unpredictable approval process” as reasons for the decline (63). Clearly, the ultimate effect of increased regulatory requirements and delays is to make innovation less attractive.

In the US, it appears that increasing delays in the approval process are partially due to a lack of trained regulatory personnel, as well as regulators’ tendency to become concerned by negative anecdotal studies. In Japan, where the approval process is even more complex, the number of experienced reviewers is even smaller and there is a large backlog of applications. It may seem curious that regulations should be multiplying even during times of reviewer staffing shortages. Some speculate that the changes may be part of a deliberate effort on the part of governments to reign in healthcare spending by reducing the approval of new medical devices and drugs. However, such a practice may in fact lead to greater social and economic costs overall. Improvements in technology ultimately lower the cost of quality healthcare. For example, since CT was first introduced, the number of CT slices per second has increased—from 1 in 1995 to 387 in 2006; meanwhile, the cost per slice has gone down, so that for the same overall price, much more information is acquired. While it is easy to add up medical expenses, it is less easy to calculate the costs (not to mention the ethical drawbacks) of not delivering efficient, high-quality medical care.

Proposals for improvements

Ironically, increasing the duration and cost of the regulatory process is only likely to increase the cost of the medical device or agent once it comes to market. To speed up the introduction of innovations into clinical practice, regulatory processes should be harmonized globally. Members of academia and industry
should encourage their governments to adopt the recommendations of the Global Harmonization Task Force, a partnership between regulatory authorities and regulated industry representatives from the European Union, the United States, Canada, Australia and Japan. In the meantime, however, industry and academia must come up with new strategies for addressing regulators’ concerns.

The development of new medical products could be streamlined through increased communication and collaboration between industry and academia. For example, the pattern in technology development has been to design a piece of equipment and then search for ways to apply it clinically—a rather inefficient process. More communication between members of academia and industry could allow technology to be developed in response to specific clinical needs (63). Fellowships enabling representatives of academia to train with industry (or vice versa) could lead to greater mutual understanding of priorities and needs, and could help improve multicenter clinical trials—say, by facilitating standardization of technology and thus standardization of institutional imaging protocols.

Efficiency in product development could also be improved by considering the whole imaging system, rather than working on different parts of it in isolation. Modern imaging requires a series of steps, including data acquisition, data reconstruction and processing, data analysis and output, and data interpretation. Different disciplines are involved in the design of the equipment used for the separate steps, including engineering, physics, and computer science. As changes in one part of the process will affect the whole, greater communication between all disciplines involved is more likely to result in advances that improve the effectiveness of the overall imaging system.

To address the need for large clinical trials, some companies have begun to conduct trials abroad. For example, General Electric has found that the total number of patients accrued at 40 sites in the US could be accrued at just three sites in India. Trial design and conduct could also be improved through more collaboration amongst industry, academia and government agencies—e.g., through organizations such as the American College of Radiology Imaging Network (ACRIN). Curricula could be redesigned to mandate training in basic and clinical research all along the educational pathway, starting with medical school. In addition, industry and academia could work together to train radiologists in the use of new imaging technology and applications even before they receive regulatory approval. This could not only improve the performance of new devices or techniques in clinical trials, but could lead to earlier, broader acceptance of the techniques and faster progress toward reimbursement.
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