Five reasons why IRs need to be part of any stroke team
Andreas Mahnken (EBIR)

Stroke is one of the leading causes of death, with an estimated 795,000 strokes per annum and 200,000 deaths annually in the US alone. It has a huge socio-economic impact. Ischaemic strokes represent roughly 80% of the total. Currently, only roughly 10% of patients receive immediate treatment, demonstrating a strong need to build up stroke units and stroke teams. Interventional radiologists with experience in neuro-imaging and intervention should be an integral part of these stroke teams and should be with the patient from the first minute. There are at least five reasons to support this claim:

1) Imaging is an extension of the clinical examination in stroke patients!
Injured but salvageable brain tissue cannot be discerned from irreversibly injured brain by means of a clinical examination. Brain imaging thus became an indispensable extension of the clinical examination. However, imaging in clinically suspected stroke is complex. Although simple unenhanced CT to exclude intracranial haemorrhage suffices to initiate intravenous lysis, diagnostic imaging has more to offer. The multimodal imaging approaches permit the detection of small infarcts and high resolution imaging of the intra- and extracranial vessels. However, images of the brain and its vascularisation need to be interpreted in the clinical context, as there are sometimes only subtle changes indicating critical pathology. Although most institutions have standardised CT and/or MR imaging protocols comprising morphologic and functional imaging aspects, there is a tendency towards individually tailoring specific components of the imaging protocol towards the clinical question. Modern stroke imaging focusses on the detection of the ischaemic penumbra, the area that can be salvaged by revascu larisation therapy. Some groups have even suggested overcoming the strict time windows for initiation of any revascu larisation therapy and tailoring any therapy on the volume of brain tissue that might be salvaged by reperfusion. With multimodal imaging being an extension of the clinical exam, presence of a radiologist is mandatory in any stroke team.

2) Interventional radiologists will optimise diagnostic algorithms!
Interventional revascularisation has become a cornerstone in the management of acute stroke. Although only 7-15% of strokes may be suited for interventional therapy, this still holds the potential for 58,000-120,000 interventions per year in the US alone. Optimal decision-making on treatment, including the planning of the actual procedure, whether intervention al, medical or surgical, is based on dedicated imaging protocols which are tailored on the individual clinical question. In fact, imaging has become an overall approach towards patient management in stroke. The IR is the single person who holds the competence in diagnostic imaging and interventional therapy and therefore is a key player in any stroke team.

3) Interventional radiologists perform best in interventional stroke therapy!
Interventional radiologists are the group with the most extensive and most regular experience in supra-aortic and intracranial interventions. In comparison with other non-radiologist groups of physicians, it has been shown that interventional (neuro)radiologists have the lowest complication rates in carotid stenting. Were similar principles to be applied to cerebral interventions, similar outcomes would be expected.

Moreover, there are a variety of different techniques and devices available. To achieve optimal results, the right device has to be selected and sufficient experience with that device is needed. Thus IRs, ideally specialised in neurointerventions, need to be part of the stroke team, as they are the ones who most safely provide interventional therapy.

4) Interventional radiologists save brain if part of the stroke team!
“Time is brain” is a well-founded and generally accepted concept in stroke therapy. There are currently time windows of 4.5 hours for i.v. lysis and up to 8 hours for mechanical revascularisation. Still, the sooner therapy is initiated the more brain tissue may be salvaged. Thus early therapy requires rapid decision making based on competent real-time image interpretation and immediate start of therapy. Both of these are key competences in stroke management, which interventional radiologists can best provide. In order to minimise delay, interventional radiologists need to be part of the stroke team starting from the first minute.

5) Radiology is a driver of the future success of any stroke team!
Actively developing diagnostic and interventional methods is a core competency of interventional radiologists. Thus, their value in a stroke team reaches beyond diagnosing stroke and treating selected patients. They are the ones providing and continuously re-inventing the basis for any stroke therapy.

Andreas Mahnken holds both MBA and MNE degrees, and has authored or co-authored more than 300 peer-reviewed journal articles and book chapters on diagnostic and interventional radiology, as well as holding several patents. He has served on the CIRSE Standards of Practice Committee and is currently part of the Vascular Division of the ESR Online Editorial Board.

Considering the huge gap between technical and clinical success in several studies on interventional thrombectomy, scientific advances (e.g. in terms of patient selection) are needed. Otherwise some promising interventional approaches may prove futile. Therefore, interventional radiologists are a driver not only in the current application, but also in developing stroke teams into the future.

In short, the management of acute stroke is rapidly developing. IRs are an asset to any stroke team as they make the difference in diagnosis and state-of-the-art therapy.

A Focus on Stroke
The Neurointerventions track in Lisbon in 2012 was hugely popular, with most rooms filled to capacity. This reflects the increasing role IR is playing in stroke management, and this year’s Neurointerventions track has been expanded accordingly, with two Foundation Course sessions, two Special Sessions, three Workshops and an exciting Hot Topic Symposium. Invited expert Andreas Mahnken lists his Top 5 reasons for IRs to get involved…

Five reasons why IRs need to be part of any stroke team

Mo 426 - 427 - 428
Building Stroke Units and Stroke Teams
Room 116
Sunday, 15 September 2013
Stroke Therapy Training – how and why CIRSE is involved

Tochi Ugbor, CIRSE Office

Interventional radiologists are playing an increasingly important role in stroke therapy. Indeed, numerous studies have already proven that IR stroke treatments such as intra-arterial thrombolysis and mechanical thrombectomy are both safe and effective when properly implemented. On examining these studies, it becomes clear that proper training is a key determining factor for the efficacy of the treatments, as practitioners require a highly advanced skill-set. CIRSE is dedicated to raising awareness of the need for better training, as well as providing IRs of all levels of expertise with valuable stroke therapy-related sessions during the annual congress.

The need for training guidelines

In a white paper published in CVIR earlier this year, IR experts Michael Lee (CIRSE President), Jim Reekers and Dierk Vorwerk express CIRSE’s position regarding stroke therapy. The document highlights the importance of adequate training, emphasising the “…direct correlation between skill level and outcome for intra-arterial stroke therapy” and foreseeing the need for more IRs to be trained to meet increased future demand. The article argues that despite the different stroke therapy guidelines that exist, “…training standards and guidelines have been less rigorously elucidated” and a “European multisociety consensus on training guidelines is highly desirable in the very near future so patients can receive high quality and safe care.”

Fundamentals of intracranial aneurysms and AVN treatment
M. Piotin (Paris/FR), C. Cognard (Toulouse/FR)

Wednesday, September 11
08:30-09:30, Room 113
WS 906
Workshop

How to perform thrombolysis and thrombectomy
A. Bernt (Augsburg/DE)

Tuesday, September 10
10:00-11:00, Room 113
FC 1001
Foundation Course

Stroke management 2: how I do it
Moderator: S. Cekirge (Ankara/TR)

Monday, September 9
08:30-09:30, Room 115
SS 203
Special Session

Preventative stroke management
Moderators: S. Cekirge (Ankara/TR), T. Engelhorn (Erlangen/DE)

Saturday, September 14
10:00-11:00, Room 111
SS 203
Special Session

Intra-arterial stroke therapy, free-of-charge, by logging into myCIRSE on www.cirse.org

7 Flodmark et al., Stroke; 2013;44:e47-e48
8 Flodmark et al., Stroke; 2013;44:e47-e48

Neurointerventions Track

Saturday, September 14
10:00-11:00, Room 111
SS 203
Special Session

Preventative stroke management
Moderators: S. Cekirge (Ankara/TR), T. Engelhorn (Erlangen/DE)

W. Kurre (Stuttgart/DE), J.M. Macho (Barcelona/ES), B. Turowski (Düsseldorf/DE), H. van Overhagen (The Hague/NL)

Tuesday, September 17
08:30-09:30, Room 111
ICS 2505
Interactive Case Session

Ischaemic stroke management – problems and solutions
T. Anderson (Stockholm/SE), T. Liebig (Cologne/DE)

14:45-16:15, Room 124
ST-Howe 2
Hands-on Workshop

Stroke therapy
Co-ordinators: J. Berkfeld (Frankfurt/DE), H. van Overhagen (The Hague/NL)

Instructors: W. Kurre (Stuttgart/DE), J. Maurer (Barcelona/ES), B. Turowski (Düsseldorf/DE), J.A. Reekers (Amsterdam/NL), J. Weber (St. Gallen/CH)

Sunday, September 15, 2013
Neurointerventions at CIRSE 2013

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COMING NEXT: ECIQ 2014, 23-26 April
When Dr. Juan Parodi, an Argentinian vascular surgeon, performed the first endovascular aneurysm repair (EVAR) in 1989, the interventional world expected that a revolution in aneurysm repair would quickly follow. Dr. Parodi initially treated Type A aneurysms with a surgical tube graft fixed only with a proximal stent. However, he quickly developed a technique to treat aneurysms extending to the aortic bifurcation and even involving the iliac arteries, utilising an aorto-uni-iliac endograft with a femoro-femoral crossover.

Fig. 1: The aorto-uni-iliac endograft first used by Dr. Parodi with femoral-femoral crossover.

Although the durability of EVAR was a concern from the start, there was the expectation that such a minimally invasive procedure compared to open abdominal aortic aneurysm repair would result in dramatically reduced morbidity and mortality. However, early registries such as EUROSTAR reported disappointing and disturbing findings, and sceptics soon appeared. In 2001, the lead article in the October edition of the British Journal of Surgery, written by the Chief Editor Dr. Jack Collins, famously declared endovascular treatment of abdominal aortic aneurysm “a failed experiment”. His criticism, based on EUROSTAR data, was focused on an unacceptable re-intervention rate after EVAR of 20% in the first year and 10% per year for at least the first four years afterward. EVAR was also 50% more expensive than open repair.

Large randomised controlled trials (RCTs) comparing EVAR to open repair for AAA did not provide reassurance for critics. The British EVAR 1 Trial and the Dutch DREAM Trial were well-designed large trials. Both trials showed a significant lower 30-day mortality with EVAR compared to open repair. However, disappoint - intimately, neither trial has demonstrated a long-term mortality benefit for EVAR with follow-up now beyond eight years. The trials conclude that “Achilles heel” recognised by Dr. Collins – a long-term re-intervention rate much higher than open repair. The excessive cost of EVAR compared to open surgery was also confirmed in these trials.

The complications requiring re-intervention in those large RCTs were predominantly due to endoleaks with device migration and graft limb occlusion also being significant issues. Since the RCTs, a number of early devices have been dis -carded and superseded by modern devices. There are large industry-sponsored device registries available to evaluate the performance of these devices. These registries have shown (as EVAR 1 and DREAM did) that EVAR provides excellent protection from aneurysm-related mortality, but at the expense of re-intervention. Although the incidence of device migration has been reduced by improved device fixation, there remain a problem, especially Type II endoleaks.

Another limitation of EVAR devices is an inability to treat the majority of AAA morphologies. This remains a challenge for modern devices. A large proportion of patients are currently being treated by EVAR with devices outside their company’s instructions for use. Infra-renal neck anatomy is the most common limitation, espe -cially in women. The reason that more patients cannot be treated by EVAR is because almost all current devices seal the aneurysm from pressure and rupture risk is by graft wall apposition at the proximal and distal sealing zones. This necessitates a parallel length of artery (typically infra-renal aortic neck and iliac artery) of good length and quality without significant thrombus, angulation or dilatation. Approximately 40% of infra-renal AAs lacks these features.

Given the persistent re-intervention rate and limited applicability of EVAR, it is not surprising that open surgical repair is still considered the “gold standard” for AAA repair. Open repair is extremely durable with a very low re-interven -tion rate and the results in the aneurysm sac being ablated and aortic side branches ligated. Conversely, the aneurysm sac is not directly treated with EVAR, leading to the risks of device migration and endoleaks, especially Type II endoleaks from uncontrolled aortic side branches arising from the aneurysm. There is much debate about the significance of Type II endoleaks, but what is beyond debate is that patients with a Type II endoleak have a much higher incidence of aneurysm sac enlargement and re-intervention. It is also clear that current treatment strategies for Type II endoleak (including intra-arterial catheter directed embolisation and percutaneous direct aneurysm sac puncture) are not particularly effective at preventing ongoing aneurysm sac growth.

Since the early days of EVAR, there have been attempts to seal the aneurysm sac at the time of the procedure, preventing complications and the need for re-intervention. In the early days of aorto-uni-iliac endografts, some centres ablated the aneurysm sac via a large sheath introduced from the contralateral groin. Sub -sequently, the most common techniques have involved pre-procedural branch artery embolisation and a catheter left in the aneurysm sac after the EVAR device has been deployed with subsequent ablation of the aneurysm sac using coils, thrombin or liquid embolic agents such as cyanoacrylate or Onyx. Unfortunately, these procedures have proved time consuming, expensive and not particularly successful.

Until recently, there has been no EVAR device that directly sealed the aneurysm sac. The Endologix Nellix device has been developed to achieve sealing of the entire aneurysm. Originally planned to consist only of two endo bags, the device has rapidly evolved to a commercial sys -tem with CE mark that involves ‘kissing’ chro -mium cobalt balloon expandable stents and surrounding endo bags (Fig. 3). The endo bags achieve aneurysm sealing by being filled by a polymer that quickly cures to the consistency of a pencil eraser. The compliant polymer filled endo bags form a cast of the aneurysm blood lumen that potentially reduces endoleak and device migration. Other potential advantages of this technique include the ability to treat a wider range of adverse proximal neck anato -mies, endoleak and device related mortality, evidence of a persistent endo -leak or device migration has been seen to date.

There is obvious potential to extend the EVAS concept beyond elective repair of AAAs. The simplicity of the EVAS procedure is attractive for the repair of ruptured AAAs and the compli -ance of endbag sealing is ideal for the parallel graft (snorkel, chimney) repair of juxta-renal aneurysms. Both procedures have already been performed with the Nellix system. There are also concepts to take this technology above the diaphragm into the thoracic aorta.

It is not surprising that other technologies are being developed to achieve aneurysm sealing. Technologies currently under development include injectable haemostatic foam implants capable of filling the aneurysm sac without endangering aortic branch arteries. These implants could be used with conventional EVAR devices to achieve aneurysm sealing. In the context of a successful endovascular aneurysm sealing (EVAS) technology is exciting and po -tentially disruptive for current AAA management. If durability can be achieved without significant secondary intervention, the current post-EVAR imaging surveillance protocols will need to be altered with major cost savings. Patients could be discharged after EVAS without the need for surveillance or secondary intervention! A revolution here has truly arrived.

Andrew Holden
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Andrew Holden is Director of Interventional Radiology at Auckland City Hospital and Associate Professor of Radiology at the University of Auckland. He is also lead Radiologist for the Auckland Hospital organ transplant programme. Dr. Holden is a committee member of ISRA (Inter -ventional Society of Australasia) and ARMS (Abdominal Radiology Society of Australia and New Zealand) and is an examiner for the RANZCR.

Dr. Holden is the author of over 60 peer-reviewed articles and three book chapters. He has been the principal investigator in 25 ‘first-in-man’ device trials and has performed over 50 live intervention -al cases broadcasts from Auckland Hospital to overseas sites such as Germany, France, Hong Kong, the USA and Australia.
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Innovating for life.
Imagine if there were no radiation required for the real-time imaging during interventional procedures. It could make us “fighter” by some kilogrammes, with no strain on the back or headaches. Unfortunately we are not there yet and we have to live as we are with our protective apparel, treat it very carefully and make sure that it will be replaced if there is any malfunction.

Is low-dose ionising radiation really dangerous for our patients and for us? After all, we only use it for noble purposes, and the dangers do not seem real but instead like the abstract result of some calculations and complex extra- polations based on several theoretical models.

Interventional radiologists are well-educated in all aspects of modern medicine, particularly imaging, but when it comes to the hardware we use daily, there is a gap in our knowledge that needs to be addressed. When buying a new car the buyer knows almost everything about it, yet basically relies on the vendor and the marketing involved. We can draw a parallel between fuel consumption of a car at the factory as opposed to what we learn when we visit the petrol station. The same could apply to radiation exposure and some other parameters published by vendors compared to what happens in “real life.” The difference is that we are usually unaware of the differences. New cars have new technology to reduce fuel consumption and to improve safety to reduce traffic accidents, while new interventional X-ray systems have technology that allows them to reduce the exposure to patients and staff, thus improving radiation safety. IRs have to learn how to master these new technologies and to use them properly, and dosimeter readings.

There are different theories regarding possible dangers of exposure to personnel. Some claim that at low doses the danger is only statistical—though the danger of constantly driving through yellow/red traffic lights on street corners is also statistical. It’s just a matter of the cumulative chances and when the damage appears.

It is extremely important to adapt our behaviour and our working culture to the powerful new X-ray machines. We have to fully understand their whole potential and be able to implement it into our daily practice.

Below-the-knee revascularisation procedures are evolving and have better outcomes than ever before. They will ultimately replace most of the distal bypasses, but are time-consuming and require prolonged fluoroscopy time, as well as an unfavourable position for the operator, close to the radiation source in the worst possible angulations of the C-Arm.

Such practice thus requires better and heavier means of personal protection. All of the above will sooner or later cause musculoskeletal problems, as well as potential and cumulative damage to our eyes.

In the era of hybrid rooms with a multidisciplinary team working shoulder-to-shoulder, we also need behavioural adaptation, as we have to orchestrate a small symphony, which comprises diverse staff members and different tools—surgical and endovascular, as well as an imaging chain with flouro or DSA, C-arm angulations, isocentric positioning of the central beam and more.

Somewhere, radiation exposure and exposure doses have become a major issue in diagnostic radiology, mostly in CT and nuclear imaging, and have been more or less neglected and often ignored in interventional radiology. Any IR has to routinely use pre-acquired CTA images, thus avoiding unnecessary DSA runs during the procedure, as well as detailed planning of each and every step of the intervention.

During recent years, CIRS has taken a very strong position in order to change this paradigm towards safe and skilled use of radiation. At the CIRS 2012 General Assembly, a number of Subcommittees were established within CIRS for permanent consultation with the Executive Committee on specific medical topics. One of the major topics is the radiation protection of patients and personnel. CIRS as a professional society has emphasised the importance of patient and particularly personnel safety. The issue of personnel safety is usually managed at national levels by relevant governmental authorities and traditionally overlooked by professional societies, which naturally promote the appealing, hi-tech part of the profession.

In an effort to further improve the society’s efforts to promote excellence in patient and personnel safety and co-ordinate CIRS’s activities in the field, a permanent Radiation Protection Subcommittee has been established and the first meeting was held during ECR 2013 in Vienna.

The Subcommittee’s core tasks are as follows:

- To provide internal and external consultation on radiation safety measures for the protection of patients and staff in interventional radiology.
- To co-ordinate all CIRS activities relating to radiation protection.
- To represent CIRS in international research consortia dealing with radiation protection.
- To evaluate documents received by CIRS dealing with radiation protection.

A dedicated group of expert IRs and medical physicists has been appointed. We believe that the activities of this group will contribute to the safe-IR practice in Europe.

CIRS also took an active part in the EU-initiated Medical Radiation Protection and Training (MEDRAPER). Following successful completion of the project, a Permanent Multidisciplinary Working Party was appointed in order to draft and maintain European learning outcome inventories for the radiation protection education, training and continuous professional development required for medical professions involved in work with ionising radiation.

Any doctor looking to improve patient and operator safety should be sure to join us today for our session on radiation dose management.

The session comprises four talks that will provide comprehensive state-of-the-art coverage of main relevant topics of radiation protection and dose management issues, including:

1. Understanding the pros and cons of the new X-ray systems in improvement of radiation safety and image quality.

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Irreversible Electroporation (IRE): Pancreatic Applications
Raj Narayanan

Pancreatic adenocarcinoma has the highest case-fatality rate of any malignancy, with a dismal overall 5-year survival rate of 5%. In the United States: 44,000 new cases of pancreatic cancer will be detected annually, with 37,400 deaths attributable to the disease. Surgical resection offers the best chance for cure, but only 10-20% of patients will qualify.

Approximately 40% of newly diagnosed patients fall in the locally advanced pancreatic carcinoma (LAPC) category, with disease which is non-metastatic, but still unresectable due to encasement of major vascular structures. Median survival remains at 6-11 months. One third of patients who initially have LAMP may become resectable after neo-adjuvant therapy. Surgery in patients with LAMP has not shown survival benefit. Intra-operative RFA and micro-wave ablation have been used with limited success, high complication rates and are associated with significant morbidity.

Chemotherapy response in pancreatic adenocarcinoma was published in 1961, utilising single-agent 5-fluorouracil (5-FU). For the next 30 years, there was no significant breakthrough in combination- or single-agent therapy, until gemcitabine became the standard of care in 1996. Combinations with Gemcitabine and Capcitabine (Cunningham et al.), and Gemcitabine with Erlotinib have been tried.

Since 2010, FOURIRINX has been used as a first-line treatment option in patients with a good performance status. A Phase 3 randomised trial comparing Gemcitabine and Folfirinox (342 subjects) was conducted by Connolly et al. [1] and demonstrated that Folfirinox was associated with a statistically and clinically significant improvement in the Median Overall Survival compared to Gemcitabine (11.1 vs. 6.8 months), but had more Grade 3 myelosuppression, febrile neutropenia, diaphoresis and sensory neuropathy.

Irreversible Electroporation is a technique that involves the use of electrodes to deliver high-voltage direct current as high as 3kV to the tumour, creating multiple holes in the cell membrane and irreversibly damaging the cell’s homeostatic mechanism, resulting in apoptotic cell death (2,3). It is cleared by the US Food and Drug Administration under the 510(k) mechanism for ablation of soft-tissue tumours, and use in the pancreas is considered to represent off-label use.

Because of its mechanism of action, tumours in contact with vessels, can be treated with IRE without compromising the vessels or creating a heat-sink effect (2,3). The preservation of vascular and ductal structures within the treatment field of IRE is hypothesised to result from the supporting connective tissue matrix, which is unaffected by this modality as a result of the lack of thermal effects (2). IRE has been studied in preclinical and clinical studies in multiple tumour types, including pancreatic cancer, and the preliminary data supports its safety and further development [4-12].

Single Centre Experience
We treated our first case of locally advanced pancreatic carcinoma in November 2010 at the University of Miami. Our initial treatment cohort consisted of 14 patients and 15 IRE treatments of the pancreas, all done percutaneously using CT guidance and under general anaesthesia. This preliminary data was presented as an abstract at the Society of Interventional Radiology in 2012 and published in the Journal of Vascular Interventional Radiology the same year (13). Two patients were down-staged for surgical resection post IRE at 4.3 and 5.3 months respectively, both obtaining negative margins post-resection. One of these cases is illustrated in Fig. 1.

Fig. 1: (a) Pre-IRE CT of patient demonstrating pancreatic carcinoma with vessel encasement (b) CT-guided percutaneous IRE (c) Pathology specimen during surgery demonstrating fat necrosis (d) Post-surgery CT follow-up

We have continued treating pancreatic cancer patients percutaneously and received IRE approval for a retrospective analysis of 30 patients who underwent 33 irreversible electroporation procedures. Three patients had two treatments each. Male to female ratio was 16:14:21 patients presented with localised disease and 9 with metastatic disease. The location of the pancreatic lesions included, head (N = 16), uncinate process (N = 2) body (N = 5), neck (N = 3), pancreatic bed (N = 4) (between the gastroduodenal and fundus and body and tail (N = 1)), surgical bed in the tail region (N = 1) encaising vessels (N = 2). Tumour size ranged from 1.2 cm to 6.8 cm, with a mean of 3.4 cm. All patients received at least one line of chemotherapy prior to IRE and some received up to five lines, and 60% of the patients also received radiotherapy.

From date of diagnosis to IRE procedure, a median of 13.9 months was calculated. Patients were down-staged for surgical resection of pancreatic cancer with vessel encasement (2,3) and a good metastatic work-up cannot be ascertained. Our median overall survival of renal cell carcinoma: a first- in -man phase I clinical report. J Vasc Interv Radiol 2012;23:142-145

Conclusion
Our initial experience with percutaneous IRE has shown that the procedure can be performed safely in carefully selected patients with the paucity of treatment options for pancreatic cancer, this presents a new treatment option. The importance of a multidisciplinary approach and a good metastatic work-up cannot be stressed enough, along with the need for a safe access for needle placement. While we have a long way to go before IRE can be incorporated into the management algorithm of the pancreatic cancer, the median overall survival data in our limited series, shows promise that will need to be further validated.

References:
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When we start PAE we should have clear in-
clusion and exclusion criteria. After undergoing patient selection, there will be unilateral
CT angiography (CTA) or pelvic magnetic resonance
angiography (MRA) before PAE. This is to exclude
prostate masses with pelvic vessels for tortuosity and atherosclerotic changes of the iliac and prostatic arteries.

A specific CT angiography protocol is applied and post-processing using maximum intensity
projections (MIP) and volume rendering with 3D reconstructions are obtained. The anatomy and atherosclerotic involvement of the iliac and prostatic arteries, the degree of calcium and stenosis of prostatic origin could by this method be known in advance, before the procedure. CTA avoids catheterisation of all other pelvic arteries and its use will ultimately reduce complications. Patients with advanced atherosclerotic involvement of the iliac and prostatic arteries are excluded on the basis of angio-CT.

With the help of angio-CT, DSA and road-
maping, the prostatic arteries are catheterised with a coaxial micro-catheter. If there is only one prostatic artery, the micro catheter should be placed before bifurcation in order to embo-
 lite both prostatic branches. The end point is the embolisation of prostatic branches, with no reflux into the arteries and opacification of the gland. If there are two prostatic arteries on one side, one should start by the anterolateral or crural branch as these are the lateral part of the prostate. So if this branch is well embo-
 lised, we should not be worried about the caudal branch. Nevertheless, it is sometimes also embolised by collateral circulation through anastomosis.

**Outcomes**

In spite of the excellent results of LIFE and its similarity to prostatic artery embolisation (PAE), the first case of PAE was performed in 2000 by DeMeritt in a patient with acute urinary retention due to BPH. In 2009, we performed the first PAE in a 76-year-
old man on acute urinary retention and bladder

Outcomes

in 2010. Carnevali et al. presented the prelimi-
ary results in two patients with acute urinary retention due to BPH successfully treated by PAE. In 2011 (10/11 patients) 91%. All patients were in acute urinary retention with bladder catheter. Patent unilateral spontaneous

in 11 of the 12 patients (90.9%). The patients did not feel any pain during or after the procedure, except one that felt pain in urinary retention before embolisation. The vesical catheter was removed 5 days after the procedure in two patients and 10 days in the remaining ones. The symptoms improved in all the patients in whom the embolisation was successfully performed (mean decrease in the IPSS of 8.2 points at 1 month and 9.3 points at 3 months). The mean prostate volume decreased from 96.3 to 74.3 ccm (22.9%) at 1 month and an additional 9.8% at 3 months. The peak urinary flow rate increased 3.8 ml/sec at 1 month and an additional 1.5 ml/sec at 3 months. At the third month patients urinated without bladder catheter with a mean IPSS of 6.33 and a peak urinary flow rate of 9.6 ml/sec.

At SR 2011, we presented the short and me-
dium-term outcomes of PAE in BPH. PAE was technically successful in 11/12 patients (98.9%) and the embolisation was bilateral in 63 and unilateral in 3. In 62 patients with clini-
success, at 12 months follow-up, all the evaluated parameters had significant clinical improvement. The remaining 4 patients improved; however, the changes were not significant, and so are considered stable in case of failure. The occurrence of a major complication, a 1.5 cm² sized blad-
ner wall ischaemia that was treated by surgical removal in two patients and 4-8 hours after the procedure, and the remain-
ing ones were discharging the morning after.

**More recent findings**

At SR 2012, Carnevali et al. presented eleven patients treated between June 2008 and November 2010. There was a technical failure (bilateral embolisation) in 75% and a failure in 25% (10/11 patients) 91%. All patients were in acute urinary retention with bladder catheter. Patients were treated spontaneously. The procedure lasted 4-25 days (mean 12.1) after catheter removal. Clinical overall improvement in LUTS at one-
year follow-up was observed by IPSS (mean 2.2) and QoL (mean 0.25). Minimum residual Volume (a teaspoon amount) was observed in 3/12 (25%) and focal bladder ischaemia in 1/12 (8.3%) procedures.

Recently we reported the short and medium-
term results of PAE in 89 patients. There were 3 technical failures (3%). PAE was bilateral in 86 patients (92%) and unilateral in 7 patients (8%). At 1 month follow-up, IPSS decreased by 10 points. QoL score decreased by 2 points, peak urinary flow increased by 38%, prostate volume decreased by 20%, post-void residual volume decreased by 30 ml and IIEF score in-
creased by 0.5 (all differences were significant, \(p<0.01\)). These changes were sustained through-
out the observation period. Seventy-eight of the 86 patients (91%) were discharged from the hospital 6-8 hours after the procedure. The remaining eight patients were discharged the following morning, 18 hours after the procedure. The bladder that each patient ranges from 2.121 to 9.766 dV cm³ (mean 3.050 dV cm³).

**Conclusion**

In conclusion, it is very important to know the prostatic arteries anatomy through previous angio-
CT or angio-MR in order to plan the pro-
cedure in advance and reduce the procedure and fluoroscopy time.

Today, more than 500 patients with BPH have been treated with PAE around the world with good and satisfactory results at both short and medium-term follow-up. In Europe, the United States and Brazil there are already several cen-
tres performing PAE. As of April 2013, we have treated over 400 patients with BPH, 17 of them with at least 3 years’ follow-up. Although we work with urologists, we have a lot of patients coming directly to us to be evaluated due to the results of the technique.

The data available in the literature are still limited and multicentric and more randomised studies are needed. With the results from two centres we can see a real benefit of PAE in selected patients with BPH.
CIRSE 2013 LIVE

The CIRSE live stream returns this year, having been an instant hit when it made its debut last year. CIRSE 2013 LIVE will be hosted on www.ecir.org and is a great way to share some of the highlights of the Annual Meeting with those who are unable to attend.

Bring your family to the Awards Ceremony via CIRSE 2013 Live and let them see you going onstage to collect your award. Or get your colleagues back home to join you for a cutting-edge lecture, so you can discuss it together when you return.

10:00-11:00
Session 102 - Upper extremity PVD

102.1 - Optimal imaging assessment for supra-aortic and upper limb arterial disease
C. Patel (Singapore)

Gore live sites:
P3 102 - Ablative therapy
Saturday 10:00-11:00

Check the live stream programming:
Saturday | Sunday | Monday | Tuesday

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It’s Simple.
At last year’s congress, it was announced that ECIO would move from a biennial event to an annual one. 2013 saw the first proper annual occurrence of ECIO, held in June rather than April, due to other fixtures in the CIRSE calendar.

Nevertheless, the meeting proved hugely popular, with over 900 participants from 60 countries making the journey to Budapest in high summer.

Despite the incredible heat outdoors, the congress centre was ideal for our needs, providing a cool and comfortable environment in which to explore the world of interventional oncology. The compact layout of the venue allowed for easy movement between sessions, booths and workshops, while still offering plenty of space and excellent facilities.

Diversity of delegates

The meeting, as always, featured an interesting mix of participants, with many representatives and speakers from other medical fields and from around the globe. This was helped in part by the ‘Bring Your Referring Physician’ programme, which offers free registration to non-radiologist colleagues and is now in its third year. Also notable were the number of young speakers presenting the research being carried out at their institutions.

Scientific highlights

Amongst the vast array of excellent lectures, workshops and discussion panels were some sessions worthy of particular attention. This year’s Honorary Lecture was delivered by Carlo Bartolozzi, a renowned researcher and educationalist from Pisa, Italy, whose career has inspired generations of Italian interventional oncologists. His lecture, Diagnosis and treatment of HCC: from guidelines to clinical practice, was a fascinating overview of the evolution of hepatic cancer therapies, as well as the current best-practice.

The ECIO meets ILCA, the World Conference on Interventional Oncology (WCIO) and our own Riccardo Lencioni, a renowned researcher and educator from Pisa, Italy, whose career has inspired generations of Italian interventional oncologists. His lecture, Diagnosis and treatment of HCC: from guidelines to clinical practice, was a fascinating overview of the evolution of hepatic cancer therapies, as well as the current best-practice.

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In a recently published paper, a panel of scientists has recommend that patients with osteoporotic vertebral compression fractures (VCF) should be treated with a procedure which reduces the impact of vertebral fractures. The recommendation reflects how I have been treating VCF patients in the past, and it is increasingly understood that both procedures induce a different impact on patients. The NICE multiple technology appraisal acknowledges that BKP may be associated with greater benefit than VP. Indeed, the SAFE trial demonstrated a 23% higher mortality risk reduction for BKP versus VP. Dr. Maneesh C Patel (UK)  You have been using the Kyphon® Cement Delivery System (CDS). What are the advantages you see with this system? I could not imagine performing a BKP or VP procedure without Kyphon® CDS gun. There are a number of benefits in using the CDS, but my primary rationale is to gain protection from radiation. The length of the connecting tube is 120 cm (this is considerably longer than other systems) which means it can keep a safe distance from the fluoroscopy unit during screening, minimizing radiation exposure. A recent publication has demonstrated that, when using the CDS gun, the radiation dose received by the operator’s finger, wrist and leg was reduced by greater than 80%, the only trial of its kind. Furthermore, the control over cement injection is significantly improved with 0.2cc’s of cement being delivered with each trigger depression. A further advantage is the ability to immediately stop the flow of cement with the safety button on the device handle which has improved the safety of cement injection and increased my confidence particularly during complex malignant vertebral fractures. It has become my cement delivery device of choice for either BKP or VP. In my practice, I tend to see older fractures which means it can be more challenging to gain height restoration. I tend to use BKP to create a safe cavity in which to inject cement in higher risk patients. However, it is accepted within studies (such as the recent FREE Surgical trial below) that one can achieve some height restoration. The FREE Surgical trial (an extension of the FREE results) demonstrated that, at 24 months, the change in kyphotic angulation was significantly improved by an average of 1.1° with BKP (vs baseline), compared to a non-significant improvement of 0.8° with NSM (p = 0.003). Importantly, within the study, this kyphotic correction improvement correlated with an improvement of Quality of Life and functioning, proving that patients with higher kyphotic restoration gain greater clinical benefits. In my opinion, there is a clear difference between the impact of different types of vertebral augmentation procedures on kyphotic correction capabilities, but there is a place for both VP and BKP depending on the patient and their fracture. In practice, I tend to see older fractures which means it can be more challenging to gain height restoration. I tend to use BKP to create a safe cavity in which to inject cement in higher risk patients. However, it is accepted within studies (such as the recent FREE Surgical trial below) that one can achieve some height restoration. The FREE Surgical trial (an extension of the FREE results) demonstrated that, at 24 months, the change in kyphotic angulation was significantly improved by an average of 1.1° with BKP (vs baseline), compared to a non-significant improvement of 0.8° with NSM (p = 0.003). Importantly, within the study, this kyphotic correction improvement correlated with an improvement of Quality of Life and functioning, proving that patients with higher kyphotic restoration gain greater clinical benefits. In my opinion, there is a clear difference between the impact of different types of vertebral augmentation procedures on kyphotic correction capabilities, but there is a place for both VP and BKP depending on the patient and their fracture.
Haemoptysis remains a severe medical condition despite advanced medical imaging techniques. It is still a life-threatening respiratory emergency and in many cases needs immediate therapy. Worldwide, the most common cause of haemoptysis remains active tuberculosis, in developed countries, the origin of haemoptysis is more often a chronic inflammatory process due to infection or conditions like cystic fibrosis, bronchiectasis, bronchogenic carcinoma or congenital heart disease [1].

In patients with massive haemoptysis, the diagnostic work-up in most cases includes contrast-enhanced chest CT alone or in combination with bronchoscopy. CT is not only able to show the bleeding area, but also to localise the exact site of bleeding in the majority of patients with haemoptysis [2].

Anatomical considerations

The anatomy of bronchial arteries is very variable in terms of origin, branching pattern and course. Bronchial arteries can arise directly from the descending aorta between the levels of thoracic vertebrae 5 and 6. Alternatively, they can arise from the thoracic aorta or arch outside vertebral 5 and 6. About 20% of the bronchial arteries arise from different thoracic or abdominal arterial branches, such as the subclavian artery, internal mammary artery or coeliac trunk. Cauldwell has defined four different branching types for bronchial arteries which arise directly from the aorta [3]. In the majority of patients, there is one single bronchial artery on the right side with distal branchings to the vessels and in general are used in sizes larger than 250 μm in order to avoid bronchial necrosis [7]. Microspheres are another possibility, as due to their hydrophilic nature and smooth spherical shape, they are less prone to clumping compared to polyvinyl alcohol particles. For bronchial artery embolisation they are used in sizes of between 500 and 700 μm [8]. In a recent study n-butyl-2-cyanoacrylate was used as liquid embolising agent in patients with bronchial artery bleeding and compared to the performance of polyvinyl alcohol particles [7]. It showed higher haemoptysis-free survival rates, without increasing complication rates. Nevertheless the use of this agent needs a high grade of experience in both delivery and especially mixing it with lipiodol in order to achieve the perfect concentration. Another exotic liquid embolisation agent for bronchial artery embolisation is ethylene vinyl alcohol polymer [9] with the only drawback being that it is relatively expensive compared to the other agents. One more possibility is the use of metal coils – detachable or not. The disadvantage of this method is that the occlusion is achieved relatively proximally in the vascular bed, which may result in recurrent haemoptysis with no possibility to do the embolisation distal to the coils.

Outcomes

The immediate clinical success rate of bronchial artery embolisation in different studies using different embolisation agents was shown to be between 73 and 100%. In the majority of studies, success rates are around 90%. Clinical recurrence rates are between 10 and 55%, with the majority around 25%. Complication rates are reported between 0 and 27%, with the majority around 10%. In case of recurrent haemoptysis, embolisation can be performed repeatedly.

In the hands of highly experienced interventional radiologists, bronchial artery bleeding embolisation is a safe and very effective method of treating life-threatening haemoptysis. Advanced knowledge of bronchial artery anatomy is necessary in order to ensure a safe and successful procedure. Contrast-enhanced multislice CT is the method of choice for detailed evaluation, and outcome in a tertiary referral hospital. Chest 1999; 132:460-464.

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CIRSE 2013
13:00-14:00
SY 1202  “MWA: the current and future state of art“
Room 115
Moderator: T. J. Vogl (Frankfurt/DE)

1202.1  “MWA: current experience and advantages“
W. Prevo (Amsterdam/NL)

1202.2  “How to achieve a good MW ablative margin (A0): tips and tricks“
B. Gonçalves (Porto/PT)

1202.3  “The science of powerful predictability“
J. Brannan (Boulder CO/US)

September
15
Sunday

Covidien - Satellite Lunch Symposium
“MWA: the current and future state of art“
Room: 115

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Interventional Oncology in Renal Cancer: Definitive cancer surgery?

David Breen

Interventional oncology (IO) continues to impact many areas of cancer care, including hepatocellular carcinoma, neuroendocrine tumour management and painful bone metastases, both to enhance standard therapies and to offer alternatives in patients declined for other treatments. Nowhere else, however, is IO more set to provide definitive, standard of care treatment than in the management of smaller volume renal tumours. This has come about through stepwise improvements in ablative technologies, image guidance and interventional techniques, but what are the relative merits of the multiple treatment options now being offered to the patient with a small (<5 cm) renal tumour?

First, we need to deal with the debate surrounding active surveillance. Many of these smaller renal cancers are indolent, yet the fact remains that we have no reliable means of predicting their behaviour. Renal cancer remains, on a per capita basis, the most lethal of all urological malignancies. 3-4 cm tumours harbour higher grade disease in 14-25% of cases [1] and the risk of metastatic progression of <4 cm disease ranges in the literature from 1 to 6% [2]. Active surveillance is predicated on the morbidity of the intervention and while, for example, no clinician would usually advocate treating a 20 mm tumour in an 87-year-old patient, many of the tumours now detected incidentally are of 25-30 mm in size and occur in otherwise fit 75-year-olds. The European Association of Urology reckons that 75% of urological tumour board, urologists have like - minded views about through stepwise improvements in ablative technologies, image guidance and interventional techniques, but what are the relative merits of the multiple treatment options now being offered to the patient with a small (<5 cm) renal tumour?

In the late-1990s – the early days of radio- frequency ablation (RFA) of renal tumours – reasonable results could be obtained for <3 cm tumours, but the poor predictability of treat- ments – largely as a result of gas obscuration – resulted in unacceptable subtotal treatment rates of 10-20%. This was borne out in 2008 by a retrospective meta-analysis of multi- institutional data which appeared to suggest that laparoscopic cryoablation (CRA) performed better than image-guided RFA both in terms of primary subtotal treatment rates and unex- pected late local recurrence [3]. This data was however plagued by case selection bias and we must be cautious in simply interpreting these findings as suggesting that CRA is better than RFA.

More recently, interventional oncologists have realised the merits of image-guided cryoabla- tion as the ‘therapeutic’ ice ball is so readily visualised using CT or MR [4, 5]. This fundamen- tally brings control and ‘scalability’ to the treat- ment and along with techniques such as retro- peritoneal fluid (or gas) hydro-dissection, means that most tumours up to 6 cm are now amenable to image-guided ablation. Of course, micro- wave brings speed of treatment to the table but, while suitable for <3 cm cortical tumours, this technique still lacks a clear and available method of true periprocedural control and monitoring. The value of multipolar irreversible electroporation (IRE) in this setting remains to be seen.

As <4 cm renal tumours increasingly represent the bulk of renal cancers encountered by the urological tumour board, urologists have like- wise been working to finesse their techniques and reduce the morbidity of partial nephrec- tomy. In particular it was realised that hiliar clamp- ing of greater than 20-30 minutes results in unacceptable deterioration of function in the operated kidney, through warm ischaemia.

‘Non-ischaemic’ techniques such as open parenchymal clamping have reduced this form of injury. Laparoscopic, ‘stapless’ partial nephrectomy and intracorporeal suturing of the pelvicalyceal system only remain reliably within the routine skill set of a limited number of centres. Extirpative surgery however pro- vides definitive proof of margins and whole specimen histology. This remains a challenge to the interventional oncology community where in situ treatment such as ablation fails to yield whole specimen histology – in a tumour notorious for grade heterogeneity – and imaging proof of the oncological outcomes is still disputed by some clinical groups.

Will there ever be a prospective, randomised controlled trial of these techniques? This author remains doubtful that in the current environ- ment there is the finance or staying power for such studies which, given the relative indolence of disease, would need to be heavily powered to show differences between the therapeutic groups. Careful registries of complications, costs and procedural outcomes would seem to be a better way forward and CIRSE, along with other bodies, is working on this.

Fig. 1a. Briskly enhancing, biopsy proven 32 mm left lower pole renal tumour.

Fig. 1b. The adjacent bowel has been displaced with contrast tinted fluid injected to the peri-renal retroperito- neum and the therapeutic ice ball can be seen subsuming the target tumour.

Fig. 1c. CT study at 10 days con- firms complete non-enhancement of the tumour, including a sub- jacent cortical margin. This is now borne out as an effective marker of tumour destruction.

References:
3. Andina DB. Jutr BC. Cancer 2006; 110:870-874
The GORE® VIATORR® TIPS Endoprosthesis is an innovative solution for TIPS. The Only FDA and CE Mark Approved Stent-Graft for TIPS
- Unsurpassed patency
- Superior radial strength
- Device flexibility
- Brilliant visibility under fluoroscopy
- Optimal configurations for TIPS applications

ENDOPROSTHESIS COMPARISON TO BARE METAL STENTS

In a randomized prospective trial, Bureau, et al., found the actuarial rates of primary patency in the GORE® VIATORR® Device group and bare metal stent group were 76% and 36%, respectively, at 2 years (p = 0.001 – log-rank test)1. In a retrospective analysis of cirrhotic patients with refractory ascites, Maleux, et al., found that TIPS using the GORE® VIATORR® Device offers better symptomatic control of the ascites at one year follow-up and a better overall survival, compared to bare metal stents2. (Figure 1)

GORE® VIATORR® TIPS Endoprosthesis Compared to Endoscopic Band Ligation (EBL)

In a randomized, controlled clinical trial with TIPS performed within 72 hours after diagnostic endoscopy and a 1-year follow up, results demonstrated an 86% actuarial survival in the early-TIPS group versus 67% in the pharmacotherapy – EBL group (p = 0.001)3. The 1-year actuarial probability of remaining free of failure to control bleeding and of variceal rebleeding was significantly higher in the early-TIPS group than in the pharmacotherapy – EBL group (97% vs. 50%; absolute risk reduction, 47 percentage points; 95% confidence interval [CI], 25 to 69; number needed to treat, 2.1 patients; 95% CI, 1.4 to 4.0)4. The conclusion was that patients with cirrhosis who were hospitalized for acute variceal bleeding and at high risk for treatment failure, the early use of TIPS was associated with significant reduction in treatment failure and in mortality.

TIPS COMPARED TO LARGE VOLUME PARACENTESIS (LVP)

Although randomized comparisons of the GORE® VIATORR® Device vs. LVP are in progress, data from bare metal stents provide evidence of the effectiveness of the TIPS procedure compared to continued LVP in acutely patients. In a meta-analysis of individual patient data, it was reported that bare metal stent – TIPS significantly improves transplant-free survival of cirrhotic patients with refractory ascites5. The cumulative probability of developing the first episode of hepatic encephalopathy (HE) was similar between the groups (p = 0.19). The average transplant-free survival at 12, 24 and 36 months of follow-up was 63.1%, 49.4%, and 38.8% per patients allocated in the BMS-TIPS group and 52.5%, 35.2% and 28.7% for patients allocated to large volume paracentesis (LVP), respectively. (Figure 2)

GORE® VIATORR® TIPS Endoprosthesis Compared to Continued Large Volume Paracentesis (LVP)

In a retrospective analysis of cirrhotic patients with refractory ascites, who were hospitalized for acute variceal bleeding and at high risk for treatment failure, the early use of TIPS was associated with significant reduction in treatment failure and in mortality.

Health Economic Benefits

Bureau et al. reported that TIPS with bare metal stents has been less cost effective than other procedures. This is mainly owing to the monitoring and the revisions required to maintain shunt patency. It has been shown that the use of covered stents could result in cost reduction because of decreased clinical re-bleaves and decreased need for shunt revisions1. TIPS is a safe intervention that reduces the need for LVP. Careful calibration allows satisfactory relief of ascites with a low incidence of HE6. It has been demonstrated that extremely low complication rates and exceptionally high patency rates can be achieved with the use of GORE® VIATORR® TIPS Endoprosthesis. In the United Kingdom, health economic data favoured TIPS with a cost of £500 per month of patient follow-up for TIPS and £3,500 per month of patient follow-up for paracentesis. Careful patient selection for this procedure has demonstrated significant health economic benefit in favour of a dedicated TIPS endoprosthesis6.

Conclusion

A large number of published data demonstrate numerous clinical advantages of GORE® VIATORR® TIPS Endoprosthesis in treatment of patients with variceal bleeding and refractory ascites. Furthermore, GORE® VIATORR® TIPS Endoprosthesis may be associated with decreased patient-care costs compared to other therapies. Considering these results, the role of GORE® VIATORR® TIPS Endoprosthesis in the management of portal hypertension should be considered. The improvement of TIPS patency by using ePTFE-covered stents is maintained over time with a decreased risk of hepatic encephalopathy and a decreased risk of death. Furthermore, data demonstrate the clinical advantage of GORE® VIATORR® TIPS Endoprosthesis in treatment of patients with variceal bleeding and refractory ascites. Finally, GORE® VIATORR® TIPS Endoprosthesis has demonstrated a decrease in associated patient-care costs. Considering these results, the role of GORE® VIATORR® TIPS Endoprosthesis in the management of portal hypertension should be considered.

Reference:
The monitoring and surveillance of vascular access are an integral part of haemodialysis patient care, as vascular access is the Achilles’ heel for a haemodialysis patient.

The Dialysis Outcome Quality Initiative Guidelines (DOQI) and the European Best Practice Guidelines (EBPG) have provided a list of techniques that can be used for the monitoring and surveillance of vascular access. The issue of adequate target-setting can be analysed by using the published works on the clinical impact of access surveillance system introductions (the timely detection of access stenosis and access patency).

The benefits of vascular access surveillance are as follows:

a) the eradication of unnoticed deterioration of delivered haemodialysis dose caused by access recirculation caused by compromised access;

b) a higher success rate in correction procedures if performed earlier in a stenosed rather than thrombosed access;

c) the general prolongation of access patency;

d) cost savings, or at least cost-effectiveness.

Monitoring strategies include physical examination (thrill, bruit, buzz and pulsations) of the vascular access to detect signs which suggest physical pathology. Access flow measurement, duplex Doppler ultrasound and direct or derived static pressure are the frequently used surveillance tools studied in the literature, with flow-measurement (QVA) being the most widely used technique.

Early detection of vascular access dysfunction can prevent not only complications related to vascular access, but also those of insufficient dialysis.

QVA is a parameter that determines the quality of vascular access, with which measurements at regular intervals can allow one to perform early interventions. Indication for a fistulography with possible intervention is recommended at access flow decrease of 25% or more over 4 months and/or at QVA below 350-400 ml/min in arteriovenous fistula (AVF) and below 400-600 ml/min in arteriovenous graft (AVG).

For reliable trend evaluation, the flow should always be measured at the same time during dialysis, preferably during the first hour. Monitoring intervals should certainly be individualised on the basis of access type (longer in AVFs than in AVGs), actual QVA value (longer for higher flows), and QVA history (shorter intervals in new access shunts, longer in known VA with stable flow).

The regularity of the intervals is often chosen according to the current status and history of vascular access. For new AVF/AVG and when decreasing flow rate once per month, a 1-month interval is also chosen after all interventions for a period of 3 months. Constant flow-measurement interval is 1 month for AVG (DOQI) and 3 months for AVF (EBPG).

All studies on vascular access monitoring and surveillance (VAMS) systems published so far have uniformly demonstrated a significant decrease in thrombosis rates and thus in thrombectomies, regardless of the access type (decrease is apparently more pronounced in AVGs than in AVFs) after introduction VAMS. This decrease is, as a rule, accompanied by a parallel increase in the number of performed prophylactic angioplasties. With the introduction of the access surveillance, the number of surgical interventions decreased from a previous 80% of all interventions to a current 10-15%, which was replaced by balloon angioplasties.

The implementation of VAMS is associated with increase in the number of interventions. VAMS entirely changed the structure of interventions; thrombectomies and surgical procedures now represent less than 10% of all interventions while over 80% are PTAs.

The guidelines of the National Kidney Foundation Kidney Disease Outcomes Quality Initiative recommend that arteriovenous vascular access undergo routine surveillance for detection and correction of stenosis. This recommendation is based on the paradigm that surveillance of access blood flow or dialysis venous pressure combined with correction of stenosis improves access outcomes. However, the quality of evidence used to support this paradigm has been widely criticised. Some authors tested the validity of the surveillance paradigm by applying World Health Organization (WHO) criteria for evaluating screening tests to a literature review of published vascular access studies. These criteria include four components: undesired condition, screening test, intervention, and desired outcome. The WHO criteria show that surveillance as currently practised fails all four components and provides little or no significant benefit, suggesting that surveillance is a false paradigm.

Duplex Doppler ultrasonography has been used repeatedly as the surveillance technique for vascular accesses at various populations. The results are also controversial.

A larger multicentre, well-organised and randomised study with hard clinical end points to evaluate the optimal surveillance strategy for both fistula and graft with adequate sample size are required.

References:
17. Podestà WD, Martin L, Loh CE, 2011 The role of vascular access surveillance in improving vascular access outcomes. Semin Dial. [Epub ahead of print]
Restaurant and Snacks at CIRSE 2013

At CIRSE 2013, you can take advantage of the Congress Centre’s coffee bar and restaurant facilities, located in the exhibition hall.

The restaurant is open Saturday to Tuesday from 11:00 to 15:00, serving a range of hot and cold Mediterranean dishes à la carte.

From soups costing as little as €3.00 to delicious mains costing €10.00, there’s sure to be something to sate your hunger!

If you prefer a quick snack or a fresh drink, the cash coffee bar will be open all day.

With a range of hot and cold beverages, as well as pastries, salads, hot-dogs and ice creams, it’s just the thing for those with a busy congress schedule.

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Visit us at CIRSE 2013 Booth 47
A new session type makes its debut today at CIRSE - the Evidence Forum. To keep pace with the latest developments in interventional medicine, Evidence Forums will see the most up-to-date research summarised and discussed in the context of relevant questions arising in current practice. Today’s Evidence Forum will examine the case for superficial femoral artery interventions, asking whether we know the optimal treatment yet.

The rising need for interventions

Peripheral arterial disease is on the rise, and is a cause for great concern. As the traditional response to the ischaemia and non-healing ulcers that arise from this disease is amputation, it can be costly in both economic and social terms. A commonly quoted figure is that somewhere in the world, a leg will be lost to diabetes every 30 seconds. Even with the wealth of IR options available and many of these patients in the USA have a primary amputation at their initial treatment - a staggering figure when one considers that recanalisation techniques can lower the amputation rate in CLI patients from 73-95% down to just 23%. An important contributor to peripheral artery disease is occlusion of the superficial femoral artery.

Today’s Evidence Forum

A range of devices and techniques are now available for revascularisation of the superficial femoral artery in peripheral arterial disease. These treatments offer great potential in reducing morbidity and restoring quality of life. However, in light of the latest studies and the volume of data now available there is a need to compare strategies carefully. This is essential if IRs are to offer an optimal and evidence-based approach for each patient.

The results of recent and ongoing investigations are well worth discussion, but comparators can be challenging if the different figuring patient groups and lesion classifications are to be taken into account. To clarify the current state of the evidence, the speakers will present an evaluation of existing data, examining short and long-term outcomes where possible and taking into consideration relevant cost implications alongside the weight of clinical evidence. A wide spectrum of endovascular options will be compared including stent technologies, where the comparative advantages of drug-eluting stents and covered stents as compared to bare metallic stents will be examined. Drug-eluting balloons, as well as atherectomy with cutting balloons, will also be discussed.

Bare Metallic Stents

The use of bare metal stents will be discussed by Prof. Stefan Müller-Hübbeck, who will ask if the received wisdom of endovascular SFA treatment still holds, or if new approaches need to be considered.

“Educated in the philosophy of restricting superficial femoral artery (SFA) stenting to bail-out, and confronted with encouraging data for new generation SFA nitinol stents for almost 10 years, a recent paradigm shift in endovascular opinion has taken place: ‘Don’t leave any metal behind!’ This statement now hangs above our daily decision-making process like a Damocles sword. The decision of when to stay with plain old balloon angioplasty (POBA), substituting angioplasty or stenting and when to switch to drug-eluting balloon, drug-eluting stent technology or combining POBA with self-expanding bare metal stents and DES for restenosis occurs.”

Cutting balloons

Dr. Mo Hamady will be discussing several treatment options, including the rationale of new ‘cutting balloons’, and how the evidence for their use is accumulating:

“Cutting balloons consist of three to four micro-surgical blades imbedded in non-compliant angioplasty balloon. The principle of cutting balloon is to minimise vessel wall trauma by inducing radial micro-incisions in the plaque prior to angioplasty. The indications for CBA include in-stent restenosis, short segment eccentric calcified plaque and graft anastomotic stenosis. CBA has been studied extensively in coronary arteries but scarce of data is available in PVD.”

“The evidence from single centre series showed good initial technical success rate with infrequent major complications. In the absence of RCT or direct comparison between CBA and other treatment modalities, it is difficult to draw a clear consensus on guidelines that identify the role and long-term results of this technique. Further studies are needed to define the short and long-term outcome, comparing CBA with other surgical and endovascular techniques, taking into account several known confounding factors.”

New stent-graft designs

The possibilites raised by stent-grafts in treating SFA lesions will be discussed by Prof. Maria Schoder, who will also evaluate the currently available devices:

“The Gore VIABAHN endoprosthesis may represent an alternative approach for endovascular treatment of long (TASC II C and D) SFA lesions. The original Gore Hemobahn endoprosthesis was introduced in Europe in 1996 and since then, innovative design enhancements were implemented. The nitinol self-expanding polyethylene fluoropolymer (PTFE) lined stent is also the only stent-graft approved by the FDA for the treatment of SFA lesions. The current generation of the Gore VIABAHN device has a heparin bioactive surface to reduce the risk of graft thrombosis and a continued proximal edge design which may improve long-term results by reducing in occurrence of proximal edge stenosis. Two recently published single-arm studies reported 12-month primary patency rates of 73% and 78% in mean treated lesion length of 19 cm. In the VASTAR trial, a prospective, randomised, multicentre study, the 12-month primary patency rate in TASC D lesions was significantly longer in the VIABAHN group compared to the bare metal stent group.”

Other technologies

Other technologies and techniques are also under investigation, and Dr. Hamady will also address the use of atherectomy devices for SFA lesions. Dr. Hamady will also discuss atherectomy devices:

“Atherectomy devices were first introduced over two decades ago. The main advantages of this modality are to avoid the use of stents in critical levels, such as the groin and the origin of the profunda femoris artery, and to reduce the neo-intimal hyperplasia. It is particularly suitable for eccentric plaques, heavily calcified lesions, long segment occlusion and femoro-popliteal bypass. Atherectomy has been assessed in small RCTs and in single or multicentre studies. The available data show high revascularisation rates, reduction in need for stenting with no significant improvement over PTA in the main outcomes of limb salvage, and mid- or long-term patency rate. Distal embolisation remains an issue of concern during the use of these devices. Combining atherectomy with drug-eluting balloons in a promising approach, which yet to be properly evaluated. The current NICE guidelines in England and Wales acknowledge the limitations of the current evidence and advise to use this technique under controlled environment or in the context of research projects.”

The session will end with a summary of the evidence by Prof. Johannes Lammer, and conclusions will be drawn by the experts in a panel discussion, considering if there is indeed a current “optimal treatment” or if further investigations and longer term trial results are needed in particular areas.

Join us to refine your knowledge of the optimum treatment for SFA, and to hear the views of the other invited speakers!

Today’s Evidence Forum: SFA - do we know the optimal treatment yet?

11:30-12:30, Room 116

Moderator: Johannes Lammer (Vienna/AIT)

Bare metallic stents

Stefan Müller-Hübbeck (Flensburg/DE)

Drug-eluting stents

Peter Gaines (Sheffield/UK)

Stent-grafts

Manja Schoder (Vienna/AIT)

Drug-eluting balloons

Gunnar Tepe (Rosenheim/DE)

Atherectomy, cutting balloons

Mohamed Hamady (London/UK)
The International Congress on Complications in Interventional Radiology (ICCIR) is a unique event in the IR calendar, giving doctors at all levels of expertise the chance to explore the difficult but necessary subject of procedural complications.

**ATMOSPHERE**
With a distinguished faculty consisting of renowned interventional radiologists, cases can be viewed through a wide variety of perspectives and experiences. The ICCIR faculty is specially selected to create an atmosphere that is sensitive and professional.

**CASE PRESENTATION**
At the core of this congress are case presentations and case reports, allowing IRs to be open and honest in discussing the complications they have faced and how these could have been best avoided or managed.

The feedback has been consistently positive, with participants appreciating the chance to improve their understanding of potential downsides through hearing about others’ experiences first-hand. A further benefit of the ICCIR is that doctors who have had difficult cases can share their stories with others who have had similar experiences.

**CASE SUBMISSION**
The ICCIR Scientific Programme Committee cordially invites all authors who wish to present a case at ICCIR 2014 to submit their abstract online by January 27, 2014. The most interesting cases will be selected and accepted for oral presentation.

**SUPPORT**
For all case presenters, registration fees will be waived and travel support of €120 will be given. Additionally, the JR Foundation is kindly offering a number of Travel Grants to case presenters once again. Submission of your application is possible from now until February 10, 2014. The first ten applicants will receive travel support of up to €500. For detailed information and eligibility criteria, as well as other congress-related information, please visit the ICCIR website at www.iccir.eu.

**DON’T MISS IT**
We are proud to announce that the ICCIR will once again provide an open forum for the discussion of complications, giving young doctors in particular the chance to learn and benefit from the experience of their elder colleagues in a structured and meaningful way. Participation is limited to 250 participants, so early registration is advised.

We look forward to seeing you in Poertschach, Austria in June 2014!

www.iccir.eu
The RapidCross™ Rapid Exchange PTA Balloon Dilatation Catheter is designed exclusively for BTK lesions. Covidien designed RapidCross™ Rapid Exchange PTA Balloon Dilatation Catheter with a unique in-line RX port and alloy support wire for optimized kink resistance. It is the only BTK balloon that has a 170 cm working length on a tapered balloon, offering improved distal access.

The balloon delivers smooth crossing with its proprietary lubricious coating, and up to two-to-three-times faster deflation than competitive products, for increased efficiency. To address the common problem of kinking with RX balloons in BTK lesions, Covidien designed RapidCross™ Rapid Exchange PTA Balloon Dilatation Catheter with a common sphere prevent aggregation within vessel lumen, providing more surface contact with vessel intima. The hydrophilic surface and media, saline, or reconstituted doxorubicin HCI allows more complete and targeted occlusion of the blood vessels with or without delivery of microspheres.

HepaSphere™ Microspheres

Merit Medical is pleased to introduce a new, smaller 30-60 micron size HepaSphere™ Microspheres. This new size gives physicians the ability to achieve more distal occlusion and to optimize embolization by more precisely matching the sphere size to the size of the targeted vasculature.

HepaSphere Microspheres’ proprietary design allows more complete and targeted occlusion of the blood vessels with or without delivery of doxorubicin HCl for the embolisation of hepatocellular carcinoma and embolisation of metastases to the liver. When packaged in their dry state, the microspheres measure 30-60 microns, but when reconstituted, they expand to 220-240 microns. HepaSphere Microspheres rapidly absorb aqueous solutions such as contrast media, saline, or reconstituted doxorubicin HCl.

HepaSphere Microspheres compress in the vessel lumen, providing more surface contact with vessel intima. The hydrophilic surface and spherical shape prevent aggregation within microcatheter lumens and in the vasculature, promoting ease and accuracy of delivery.

Covidien is celebrating five years in safe microwave ablation with the anticipated release of the Empirin™ advanced ablation system.

Learn more about the “Science of Powerful Predictability” at the Covidien Learning center #2 and at the lunch symposium on Sunday at 13:00 in room 115.

EPIQ brings expanded performance to the Interventional Suite

EPIQ, the new premium ultrasound from Philips, introduces solutions that improve diagnostic confidence, speed decision making and improve connectivity. With EPIQ multi-modality image retrieval is available at the push of a button on the system, without the use of an external reading station. The system’s fully integrated fusion capabilities feature streamlined workflows to allow clinicians to achieve fast and effective fusion of US/MR/CT/PET with live ultrasound. Additionally, EPIQ features advanced needle navigation for challenging interventional cases, such as hard to see patients or for patients who cannot be moved. Such solutions to improve connectivity and increase speed of fusion have the potential to improve workflow and clinical confidence, even in challenging diagnostic cases.

MULTICORE® provides an optimised needle visualization under ultrasound guided biopsy procedures. By the nature of its constituent material it functions at any angle of entry into the body in relationship to the generation of sound waves by the ultrasound transducer. Thanks to its perfect smoothness, avoids any risk of seeding of malignant cells along the needle’s path from the patient’s body out.

Specimens provided through MULTICORE® are particularly abundant and allow a quick, safe and easy biopsy procedure, either performed manually or through the most common imaging guiding systems, such as CT, US, MRI.
New Product Launches

**TERUMO**

**TERUMO Europe N.V. announces the launch of MicroThermX™ Microwave Ablation System**

MicroThermX™ is a powerful, reliable and user-friendly microwave system. Microwaves technology treats tumors faster than Radiofrequency Ablation and there is less impact by heat sink effect.

MicroThermX™ is simple to use; its small and lightweight generator can easily be moved and stored. Its lightweight and flexible cables allow an easy antenna manipulation. When using multiple probes, its synchronous wave alignment technology allows flexibility in antenna placement without any risk of skin burn and creates a consistent area of necrosis.

MicroThermX™ is a CE marked and FDA approved device which is produced in the US by BSD Medical Corporation and boasts more than 1,000 successful procedures.

TERUMO was a pioneer in Interventional Oncology with the introduction of Drug-Eluting Beads technology in Europe and it aims to indorse its support to the Interventional Oncology community by offering a complete solution for Interventional Oncology.

**VIDACARE**

**Hard Bone Lesions Made Easy, with powered bone biopsies using OnControl’s new coaxial needle**

The OnControl® powered bone access system, using Vidacare’s handheld driver platform combined with procedure-specific needle sets, represents the first major advance to bone lesion biopsy solutions since 2009. Our bone lesion biopsy solutions have provided interventional radiologists a faster, more reliable tool for accessing dense and hard-to-reach bone lesions. Introduced in 2012, the coaxial biopsy tray is specifically designed for multiple bone biopsy from a single cortex penetration, and also remains clearly visible through imaging for precise placement.

For more information on OnControl bone lesion biopsy solutions, see us at booth #50, visit Vidacare.com/OnControl or email us at oncontrol.international@vidacare.com

**CIRSE 2013 Party**

Tuesday, September 17
Doors open at 20:00
MNAC, Barcelona

Cocktails with outstanding views over Barcelona! Free visit of the unique MNAC Romanesque Art Collection®
Delicious food! Great live music!

The last evening of CIRSE 2013 will be one to remember!
Get your CIRSE 2013 Party ticket now!

Dinner and party ticket: €90 per person
Party only: €25 per person

To secure your ticket, visit the Kuoni booth in the entrance hall!

CIRSE supports compliance with ethical standards. Therefore, CIRSE emphasises that the present offer made by KUONI Destination Management is directed to participants of CIRSE 2013 and recommends that the participants who want to accept the present offer shall bear any and all costs in this context themselves.

Kindly note that entrance to the CIRSE 2013 Party (dinner and/or party) is not included in the CIRSE 2013 registration fee!

**EDUCATION IN INTERVENTIONAL RADIOLOGY**

**ESIRonline**

Did you know that ESIRonline contains more than 6,100 presentations from all areas of interventional radiology?

The CIRSE 2013 presentations will be added very soon – keep your eyes peeled!

CIRSE members benefit from year-round access to the complete lecture database on ESIRonline.

www.esir.org ... Interventional Radiology at your fingertips
Germany Sets New Standards in IR Quality Management

Petra Mann, CIRSE Office

It has been a few years since the German Society of Interventional Radiology (DeGIR) implemented its ground-breaking quality management programme. The objective was to maintain and improve interventional radiology’s high standard of care and professional status across the country.

The DeGIR quality management programme addresses two key dimensions of medical professionalism – certification of interventional radiology (IR) specialisation and procedure data collection and monitoring.

Specialisation and Certification

DeGIR’s certification in IR is a structured, educational programme that encompasses a basic and an advanced level. It is compatible with similar, Europe-wide programmes (European Board of Interventional Radiology – EBIR) and is fully supported by the German Society for Radiology’s Academy for Continued Education. Thanks to its high standards and official endorsements, the DeGIR certificate guarantees holders recognition of their knowledge, skills and competencies in IR.

DeGIR’s educational programme (in full support of EBIR) is complemented by qualification guidelines drafted and published in co-operation with the German Radiology Society (DRG).

Data Collection and Monitoring

In order to provide continuous quality assurance in IR, a state-of-the-art system was developed by DeGIR which allows interventional radiologists to collect and record data pertaining to procedures they have performed throughout their careers.

This unique online system allows its members to enter an array of procedural data, including quality parameters and treatment outcomes. Users can enter data in real time, evaluate their procedures according to quality assurance parameters and compare them anonymously to all available data.

DeGIR’s dedication to this ground-breaking system is shown by the team of specialists continuously reviewing and adapting it to new procedures. This commitment is certainly paying off, with almost 200 hospital and institutes participating in the programme. In 2011, more than 80,000 procedures were entered, making it the most comprehensive collection of IR quality assurance data worldwide.

Furthermore, registration with and regular use of the system are strict requirements of the DeGIR certification.

Committed to Excellence

To date, there are more than 900 DeGIR certified interventional radiologists and 190 certified IR instructors from 120 educational centres in Germany.

The benefits that such a programme brings to the patient cannot be denied. It has also proven to be an important tool in demonstrating the high standard of IR procedures vis-à-vis more traditional methods and is highly instrumental in discussions on the distribution of resources among the various medical disciplines – which naturally requires supporting data.

Registries in Interventional Radiology

Robert Bouter, CIRSE Office

Medical case-series studies1, more commonly known as registries, are an important and often overlooked tool in bolstering the evidence base of medical interventions to help improve the quality of service. Registries involve collecting crucial data points of procedures (duration, success, complications, etc) performed in multiple centres.

Unlike other forms of clinical research, registries do not rely on a randomised sample of patients and the inclusion criteria allow for a wider patient sample.

This straightforward methodology is the greatest advantage of a registry, as Jim A. Reekers, Professor of Interventional Radiology at the University of Amsterdam points out, “registries give a grossly mode overview of the efficacy, safety, complications and frequency of new procedures. Registry data can also assist in setting up more sophisticated trial studies, helping calibrate how many patients are needed to produce good scientific evidence.”

Naturally, registries are not without their challenges, as Prof. Reekers explains, “registries do, however, have an uncontrolled voluntary inclusion and are therefore particularly susceptible to selection bias and must be interpreted cautiously.”

Owing to their manageable and effective methodology and the relatively modest setting-up costs, registries are readily applicable enquiries that help better understand and ultimately fine-tune the provision of interventional radiology services.

For examples of IR registries, please refer to www.cirse.org


High Volumes, Good Outcomes

Ciara Madden, CIRSE Office

For many years now, it has been accepted, and indeed proven, that a strong link between high-volume medical centres and good patient outcomes exists. The first study into this link appeared in 1979, when Dr. Harold Luft of the University of California at San Francisco published a paper in the New England Journal of Medicine, showing 25–41% fewer patient deaths in hospitals performing 200 or more surgical procedures a year, compared with lower volume hospitals. Since then, there have been many such studies conducted, mostly in the surgical field2, and all appear to show favourable outcomes in high-volume centres, particularly for rare or difficult cases.

The mechanisms governing such an effect have yet to be fully established, but several possible factors have been identified:

- Regular experience means a hospital and its staff are more primed to deal with such cases, and more familiar with the options and therapies.
- High-volume centres have better resources, such as imaging and diagnostic equipment, which may result in more accurate therapy delivery.
- Surgeon experience may be the underlying cause, and high-volume centres are more likely to provide surgeons with regular opportunities to practice.
- Perhaps good outcomes attract high volumes of patients, and not the other way around.

Whatever the underlying cause, it is clear that high volume, specialised centres can often offer better outcomes, and many medical centres in Europe already reflect this trend, with specialised regional centres widely offered for stroke, cancers and paediatrics, to name but a few.

The Case for Interventional Radiology

There are less data available in the case of interventional radiology (IR), although data on endovascular treatment for stroke do show similar outcomes3. Moreover, it seems logical that interventional radiologists who regularly treat particular pathologies or emergencies will become more adept at treating those cases. It is already the case that interventional radiologists who work in hospitals specialising in trauma, cancer or stroke are emerging as experts in these fields, and are in a position to train and educate their colleagues.

In the same way, centralising IR care, especially for more complex and challenging cases, could greatly contribute to enhanced patient safety – a structure best arranged amongst hospital management, interventional radiologists and medical insurance companies. Given the evidence that exists, hospital administrators would be well advised to capitalise on the resources already available to them. By ensuring that proper referral pathways and collaborative structures exist, hospital managers can ensure that their IR staff treats a regular stream of patients, keeping their hard-won skills up to date and staying abreast of new therapies and technologies – and most importantly, improving the outcomes for patients.

1 Vernooy et al. Specialised and high-volume care leads to better outcomes of ovarian cancer treatment in the Netherlands, University Medical Centre Utrecht


3 Speedier treatment and better outcomes for high-volume stroke centres, Journal of Neuro-interventional Surgery, 9th May 2012

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If you have any questions about this publication, please contact us at madden@cirse.org.

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These articles were originally published in Intervention IQ magazine. For further information on a range of IR issues, please visit www.iqonline.eu, or visit our IQ Lounge here at CIRSE.
Innovation at your fingertips

Intervention IQ is a centre for information and tools promoting IR to healthcare professionals. Available across all digital formats, IQ houses a wealth of articles and interviews dealing with an array of conditions that affect our society today, and healthcare systems as a result. Like IR itself, IQ is innovative and dynamic, having recently been digitally re-launched.

Intervention IQ app now available

IQ is building a community of healthcare professionals who are interested in IR. The Intervention IQ app, available from the Apple Store, Google Play and Google Currents, means you can follow the latest content and access the IQ archives at your convenience.

Free unlimited downloads

Information-sharing is vital for promoting awareness and information on IR. With this in mind, IQ launched the Download Centre, allowing you free and unlimited use of select content for your own meetings and presentations. Experience what’s on offer by visiting the IQ Lounge!

Connect with IQ

IQ strengthens its links with the IR community via social media as well as through the IQ blog, which is designed to meet the demand for a user-friendly and comprehensive resource for promoting awareness of IR.

Find out more at the IQ Lounge!