Limited Assurance Report

COCIR European Coordination Committee of the Radiological, Eletromedical and Healthcare (IT) Industry, SRI Steering Committee Secretariat
Brussels

Review of the SRI Status Report 2016

Contract: 0.0834930.001
Independent Practitioner’s Limited Assurance Report

To the European Coordination Committee of the Radiological, Electromedical and Healthcare IT Industry (“COCIR”), the SRI Steering Committee Secretariat of the Self-Regulatory Initiative (“SRI”) for Medical Imaging Equipment under the Ecodesign Directive, Brussels.

We have been engaged to perform a limited assurance engagement on selected data marked with ✓ in the Status Report of the SRI for Medical Imaging Equipment under the Ecodesign Directive (hereafter the “SRI Status Report”), Brussels, for the year 2016 in the context of the reporting requirements of the SRI.

Management’s Responsibility

The SRI Steering Committee Secretariat of the SRI is responsible for the preparation and presentation of the SRI Status Report in accordance with the six step methodology as stated in the SRI V1 (for ultrasound imaging equipment) and SRI V3 (for MRI, CT and future modalities) (hereafter the “SRI six step methodology”):

- Step 1: Gather baseline data
- Step 2: Prioritization and selection of next modality
- Step 3: Identification of significant environmental aspect(s) for the selected modality
- Step 4: Derive environmental targets and objectives for the selected modality
- Step 5: Implementation into company processes
- Step 6: Monitoring and reporting

and for the selection of the information to be assessed.

This responsibility includes the selection and application of appropriate methods to prepare the SRI Status Report as well as the use of assumptions and estimates for individual SRI disclosures which are reasonable in the circumstances. Furthermore, the responsibility includes designing, implementing and maintaining systems and processes relevant for the preparation of the SRI Status Report, which is free of material misstatements due to intentional or unintentional errors.

Audit Firm’s Independence and Quality Control

We have complied with the German professional provisions regarding independence as well as other ethical requirements.

The audit firm applies the national legal requirements and professional standards – in particular the Professional Code for German Public Auditors and German Chartered Auditors (“Berufssetzung für Wirtschaftsprüfer und vereidigte Buchprüfer”: “BS WP/vBP”) as well as the Standard on Quality Control 1 published by the Institut der Wirtschaftsprüfer (Institute of Public Auditors in Germany; IDW): Requirements to quality control for audit firms (IDW Qualitätssicherungsstandard 1: Anforderungen an die Qualitätssicherung in der Wirtschaftsprüferpraxis - IDW QS 1) – and accordingly maintains a comprehensive system of quality control including documented policies and procedures regarding compliance with ethical requirements, professional standards and applicable legal and regulatory requirements.
Practitioner’s Responsibility

Our responsibility is to express a limited assurance conclusion on the data marked with ✓ in the SRI Status Report based on our work performed.

Within the scope of our engagement we did not perform a substantive audit on external sources of information or expert opinions, used to prepare and referred to in the SRI Status Report.

We conducted our work in accordance with the International Standard on Assurance Engagements (ISAE) 3000 (Revised): “Assurance Engagements other than Audits or Reviews of Historical Financial Information” published by IAASB. This Standard requires that we plan and perform the assurance engagement to obtain limited assurance whether any matters have come to our attention that cause us to believe that the data marked with ✓ in the SRI Status Report has not been prepared, in all material respects, in accordance with the SRI six step methodology.

In a limited assurance engagement the evidence-gathering procedures are more limited than for a reasonable assurance engagement and therefore significantly less assurance is obtained than in a reasonable assurance engagement. The procedures selected depend on the practitioner’s judgment. This includes the assessment of the risks of material misstatements of the data marked with ✓ in the SRI Status Report with regard to the above mentioned SRI six step methodology.

Within the scope of our work we performed amongst others the following procedures:

- Inquiries of personnel at the SRI Steering Committee Secretariat (COCIR) responsible for the preparation of the SRI Status Report regarding the process to prepare the report and the underlying internal control system;
- Inspection and sample testing of the systems and process documentation for collection, analysis and aggregation of the selected data;
- Recalculation of the aggregation and KPIs calculation for selected data;
- Analytical procedures on selected data;
- Inspection of documents regarding the description of the SRI six step methodology (SRI V3) and its application to MRI as detailed in ‘Magnetic resonance Equipment (MRI) - Study on the potential for environmental improvement by the aspect of energy efficiency’ and ‘Magnetic Resonance – Measurement of energy consumption’.

Conclusion

Based on our limited assurance engagement, nothing has come to our attention that causes us to believe that the data marked with ✓ in the SRI Status Report of the SRI for Medical Imaging Equipment under the Ecodesign Directive for the year 2016 has not been prepared, in all material respects, in accordance with the SRI six step methodology.

Emphasis of Matter – Recommendations

Without qualifying our conclusion above, we make the following recommendations for the further development of the reporting of the Self-Regulatory Initiative for Medical Imaging Equipment under the Ecodesign Directive:

- Further improvement of effectiveness and documentation of controls
Restriction on Use and Distribution

We issue this report on the basis of the engagement agreed with COCIR. The assurance engagement has been performed for purposes of the SRI for Medical Imaging Equipment under the Ecodesign Directive and is solely intended to inform SRI for Medical Imaging Equipment under the Ecodesign Directive about the results of the assurance engagement. The report is not intended for any third parties to base any (financial) decision thereon. Our responsibility lies only with COCIR. We do not assume any responsibility towards third parties.

Munich, 7 September 2017

PricewaterhouseCoopers GmbH
Wirtschaftsprüfungsgesellschaft

Hendrik Fink                    ppa. Annette Daschner
Wirtschaftsprüfer             (German Public Auditor)
Index to appendices

I  Self-Regulatory Initiative („SRI“) Status Report 2016 .......................................................... 1

General Engagement Terms for Wirtschaftsprüfer and Wirtschaftsprüfungsgesellschaften
[German Public Auditors and Public Audit Firms] as of January 1, 2017
SELF-REGULATORY INITIATIVE FOR MEDICAL IMAGING EQUIPMENT
STATUS REPORT 2016

SEPTEMBER 2017
REPORT OVERVIEW

The Status Report on the Self-Regulatory Initiative for medical imaging devices, published annually by the SRI Steering Committee presents information on the SRI, the developments and concepts, and the results achieved by participating companies.

This SRI Status Report 2016 consists of four main parts and eight appendixes

Part 1 offers a general introduction to the self-regulatory initiative, describing the development of the methodology from the first proposal in 2009 (SRIv1), to the second version (SRIv2) submitted to the European Commission and the Consultation Forum early in 2012 and the present third version (SRIv3).

Part 2 lists news and results of the work of the Steering Committee and the Expert Groups in 2013.

Part 3 explains in brief the content of the six steps of the SRIv2 methodology. More details on the methodology are available in the SRIv2 documentation (www.cocir.org).

Part 4 shows the results achieved from 2011 to 2016 by participating companies for all the modalities under scope: ultrasound, magnetic resonance, computed tomography, X-ray and nuclear imaging.

Appendix I summarizes the ultrasound pilot project and briefly present the SRI V1 methodology.

Appendix II displays the results of the SRIv2 methodology applied in 2010 to all the modalities in scope of the Self-Regulatory Initiative. Step 1 and step 2 of the methodology allowed to define a Priority List based on LCA data provided by companies.

Appendix III presents the results of the application of the methodology to magnetic resonance imaging equipment that have been identified as priority one for their environmental impact. The "MRI measurement of the energy consumption" methodology is also briefly introduced.

Appendix IV summarizes the findings of the 2013 project on computed tomography equipment and the application of the SRIv3 methodology.

Appendix V presents the findings of the application of the methodology to X-ray medical equipment.

Appendix VI introduces the nuclear imaging modality, the findings and the decisions of the SRI Steering Committee.

Appendix VII presents the new concept methodology to measure the circularity of the economy of the Medical Imaging Devices Sector. This new project has been launched as the SRI Steering Committee realized that contributing to circular economy can deliver more environmental benefits than improving energy efficiency only.

Appendix VIII shows the SRI list of relevant information that an Environmental Products Declarations (EPD) should contain.

Data and figures marked with the ✓ (green) logo have been in the scope of the PwC review 2016 (see page 82 for additional information). Data and figures market with the ✓ (black) logo were in the scope of previous PwC reviews (2010, 2011, 2012, 2013,2014 and 2015)
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GENERAL INFORMATION ABOUT COCIR

Founded as a non-profit trade association in 1959, COCIR represents the Radiological, Electromedical and Healthcare IT industry in Europe. As such, COCIR members play a driving role in developing the future of healthcare both in Europe and worldwide. COCIR is committed to support its members and communicate with its partners in Europe and beyond on issues which affect the medical technology sector and the health of EU citizens. COCIR also works with various organisations promoting harmonised international standards and regulatory control that respects the quality and effectiveness of medical equipment and healthcare IT systems without compromising the safety of patients and users. COCIR encourages the use of advanced technology to support healthcare delivery worldwide. Key objectives are to include and to promote free worldwide trade of medical equipment and to maintain the competitiveness of the European health sector.

COCIR ENVIRONMENTAL HEALTH AND SAFETY COMMITTEE (EHS)

Founded in 2000 the COCIR EHS has taken several initiatives in the environmental domain introducing ecodesign initiatives in different ways:

- **2002 - 2007**, in the field of **International Standardisation**: COCIR member companies contributed to the development of an internationally-recognized standard integrating ecodesign into the product design and development process for all electromedical equipment. International Electro-technical Commission (IEC) published the International Standard IEC 60601-1-9
- **In 2006**, in the field of **Integrated Product Policy**, as advocated by the EU’s Action Plan on Sustainable Consumption and Production and Sustainable Industrial Policy (SCP/SIP), COCIR member companies participated in an Integrated Product Policy project with the Hamburg authorities and hospitals.
- Keeping up with the latest inventions in medical technology often involves replacing equipment in medical practice before it reaches the end of its useful life. COCIR published in 2007 a version 1 of **Good Refurbishment Process (GRP)** describing in 5 steps how manufacturers can effectively refurbish equipment to ensure quality, safety and effectiveness of medical imaging equipment.
- **COCIR published in 2008 a guide on REACH requirements for component suppliers and equipment manufacturers**
- **In 2008**, COCIR launched a **web-based database for substances declarations under REACH, RoHS, Batteries and Packaging directives called BOMcheck**. This centralized open-access database provides a cost-effective approach for manufacturers to work with their suppliers to reduce hazardous substances in products.
- **In 2014**, considering the importance of environment as a horizontal issue, the scope of the Committee was broadened, and now covers also health and safety.

COCIR.org

More detailed information on COCIR initiative in environmental domain could be found on COCIR website [www.cocir.org](http://www.cocir.org).

Footnotes:
1. For more information: [www.cocir.org](http://www.cocir.org).
PARTICIPATING COMPANIES

The following companies participate in the SRI for Medical Imaging Equipment for the modalities they market in EU.

**Table 1:** Participating companies in 2016

<table>
<thead>
<tr>
<th></th>
<th>Magnetic Resonance MRI</th>
<th>Computed Tomography CT</th>
<th>Nuclear Medicine NM</th>
<th>Radiology X-RAY</th>
<th>Ultrasound US</th>
<th>Radiation Therapy(^2) RT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elekta</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fujifilm</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>GE Healthcare</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hitachi Medical Systems Europe</td>
<td>(\checkmark)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IBA - Ion Beam Applications</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(\checkmark)</td>
</tr>
<tr>
<td>Philips Healthcare</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
</tr>
<tr>
<td>Samsung Europe</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(\checkmark)</td>
</tr>
<tr>
<td>Siemens Healthcare</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td>(\checkmark)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toshiba Medical Systems Europe</td>
<td>(\checkmark)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agfa Healthcare(^4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^2\) Radiation Therapy (RT) has been added in 2015 to the table as the new “circular economy” project (see Appendix 7) covers all modalities. Elekta and IBA have been moved to RT as this better represent their activities.

\(^4\) Agfa Healthcare joined the SRI in 2015 following the decision to investigate “circular economy”
1. GENERAL INTRODUCTION TO THE SRI FOR MEDICAL IMAGING EQUIPMENT

The Energy Related Products (Ecodesign) Directive, 2009/125/EC, enables the European Commission (EC) to set ecodesign requirements through new regulations for any group of products which uses energy. In 2007, Medical Devices were identified as a “Priority A” product group by the EC for future regulation. To avoid adverse business impacts (unnecessary costs and loss of flexibility in product design), COCIR reached a consensus with the EC to develop an alternative approach allowed under the Ecodesign Directive Annex VIII (Self-Regulatory Initiative for an industry sector).

1.1. SELF-REGULATORY INITIATIVE V1 - 2009

During the EC Consultation Forum meeting on 28 May 2008 COCIR presented its first proposal for an industry-led Self-Regulatory Initiative. The EC welcomed this alternative approach as it could achieve the same overall objective as an implementing regulation but would avoid potential negative business impact. In particular, the EC emphasised that “regulation would risk hampering innovation in the medical equipment sector, where technology evolves rapidly”.

Based on this positive feedback, COCIR decided in September 2008 to establish the SRI Steering Committee (hereafter: SRI SC) in order to further develop this Initiative and develop the methodology.

- In May 2009 the developed methodology (hereafter: SRI) was applied to ultrasound products in a pilot project to gather experience from practical implementation.
- In November 2009 the methodology and the pilot project on ultrasound products was presented to the Consultation Forum. Comments were gathered for further improvements.
- In 2010 COCIR and the Steering Committee worked on the SRI v1 to update and improve it taking into account the comments received from the Members of the Consultation Forum, the EC and the lessons learnt from the ultrasound pilot project.

1.2. SELF-REGULATORY INITIATIVE V2 - 2012

The result of the thorough analysis and review is the SRIv2 6 step methodology which the Steering Committee considers to address all concerns. The SRI has been submitted to the European Commission and Consultation Forum in January 2012 and the Consultation phase closed on 5th March 2012. The SRI has been officially acknowledged by the Commission in November 2012.

1.3. SELF REGULATORY INITIATIVE V3 - 2013

In 2013 the SRI Steering Committee reviewed the SRIv2 Methodology to integrate received comments and the new findings and methodologies developed during the MRI project in 2012. The methodological aspects have been separated from the commitments of signatories. Two papers are now available for download on the COCIR website:

1. COCIR Self-Regulatory Initiative
2. COCIR Self-Regulatory Initiative Methodology V3 (SRIv3)

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5 Article 18 of the Ecodesign Directive (2009/125/EC) establishes a “Consultation Forum” (CF) which allows stakeholders to be informed and consulted on the implementation of the Directive. The Forum is limited to 60 members, including representatives of EU Member State, 3 representatives of EEA Member States, 30 stakeholders (Stakeholders have been selected by the EC following an open call for interest).
2. NEWS AND DEVELOPMENTS 2016

During 2016 the SRI SC worked to evaluate the feasibility of the data collection needed to provide the inputs to the concept methodology to measure the circularity of the medical imaging business model in EU. On the basis of the results of the evaluation, the SRI SC launched a campaign to collect already available data, with the aim of calculating sub-indexes.

2.1. 5TH ANNUAL FORUM MEETING ON 11 APRIL 2016

During the 5th Annual Forum of the COCIR SRI on 12 April 2016, the SRI Steering Committee presented the results of the work performed in 2015. In particular the discussion focused on the package adopted by the EC on Circular Economy and how such concept can be introduced in the SRI methodology. The initial methodology presented during the meeting is now included in this report in Appendix VII.

The event was open to Members of the Consultation Forum, Stakeholders and Interested Parties. The documents presented and discussed during the meeting are now available at the COCIR website.

2.2. ASSESSING ENERGY SAVINGS FOR MRI

After 4 years of measuring the energy consumption of MRI equipment placed on the market, the SRI SC introduced an estimation of the total energy saved by the initiative from 2011 to 2017 in chapter 4.8.1. While previously energy savings where estimated on the basis of the projected Business as Usual Scenario and the Baseline 2011, now the savings are calculated on real data.
3. THE SRI METHODOLOGY

The SRI methodology is the process participating companies shall follow to set ecodesign targets for their products and ensure that they are achieved.

The purpose of the SRI methodology is to:
- Provide a transparent and continuous process to control the application of ecodesign targets while protecting company confidential information.
- Set a priority sequence for the equipment evaluation.
- Identify top environmental aspects.
- Set environmental targets.
- Systematically engage stakeholders.
- Monitor and report progress.

The SRI methodology is based on six key steps. The iterative process allows the member companies to offer their confidential internal life cycle assessment data for each modality, life cycle stage and aspect selection processes. This provides a comprehensive analysis which any one single approach could not achieve.

3.1. SCOPES

The SRI methodology applies to the following imaging equipment:
- Magnetic Resonance Imaging equipment (MRI)
- Computed Tomography (CT)
- Nuclear Medicine (NM)
- X-ray
- Ultrasound (US)
- Therapy equipment (under discussion)

Every year at least one new modality is selected until all modalities in scope have been chosen.

After a modality has been selected (via the priority list resulting from Step 1 and 2), it will remain in the continuous improvement circle (Steps 3 to 6). This means that once a modality has been selected and the industry has achieved the target to minimize the aspect with the highest environmental impact and potential for improvement, another assessment of the most significant aspects will be done.

3.2. STEP 1: GATHER BASELINE DATA

The purpose of the first step is to establish a baseline for prioritization and improvement. The method of the first step is to collect Member Company Life Cycle Assessment (LCA) data on products in scope of the SRI.

The provided information in Step 1 contains typical LCA data per modality and company, including the defined use scenarios, functional units, impact categories and intended use, the percentage of life cycle contribution, the individual environmental load per modality related to the corresponding LCA method, as well as a plausibility check based on expert expectations on the future capacity for innovation of the modality.

3.3. STEP 2: PRIORITIZATION AND SELECTION OF NEXT MODALITY

The purpose of the second step is to analyse the baseline data in order to prioritize the modalities for evaluation. The method of the second step is to rank the environmental load and other data from each modality.

3.4. STEP 3: IDENTIFICATION OF SIGNIFICANT ENVIRONMENTAL ASPECT(S)
FOR THE SELECTED MODALITY

The purpose of the third step is to analyse and prioritize the three most significant environmental aspects contributing to the total environmental impact of the selected modality. The method of the third step is to rank the data on the top three environmental aspects, including the percentage of their contribution to the total life cycle.

3.5. STEP 4: DERIVE ENVIRONMENTAL TARGETS AND OBJECTIVES FOR THE SELECTED MODALITY

The purpose of the fourth step is to develop ecodesign targets. First the selected modality is defined (system boundaries, categories, parameters, protocols, use scenarios, etc.) and then a measurement methodology for the chosen environmental aspect (e.g. energy consumption) is developed. The methodology allows companies to measure their equipment.

The SC Secretariat uses the above mentioned collected values and the results of the study on improvement potential (when appropriate) to calculate the target scenarios:

- Best available technology as Baseline (BAT)
- Business as usual (BAU)
- Best not yet available technology (BnAT)
- Beyond Business as usual (Beyond BAU)

Based on these scenarios, the SRI SC decides on a feasible industry reduction target. Before it is integrated into the companies’ design targets, the industry target is proposed to the European Commission for discussion.

The results of this step are two types of targets:

- **Individual company targets:** These are absolute improvement targets which are calculated on the basis of declared values and expected sales at the end of the innovation cycle. Company targets are communicated by the SRI SC Secretariat to each company in a confidential way. It remains at the company’s own discretion to publish this individual target.

- **Industry target:** This target represents the market fleet average the Industry is committed to achieve in a period of time equal to the innovation cycle.

3.6. STEP 5: IMPLEMENTATION INTO COMPANY PROCESSES

The purpose of the fifth step is the integration of the ecodesign target(s) into the internal company processes. The translation of the target(s) into the company internal product design process remains up to the individual member.

3.7. STEP 6: MONITORING AND REPORTING

The purpose of the sixth step is to monitor and report on the achievement of the industry target. The method of the sixth step is to calculate the averages of the annually reported impact values and comparison to the Baseline. The status of each modality and the progress that the companies are making is annually reported to the Stakeholders with this SRI Status Report.
4. ACHIEVEMENTS AND DEVELOPMENTS PER YEAR

2011 -

4.1. MAGNETIC RESONANCE
In 2011 the SRI Steering Committee officially launched the eco-design target for MRI after the finalization of:
- The MRI energy consumption measurement methodology
- The study on improvement potential
- The ecodesign target for 2017
- The data collection for 2010 and 2011

2012 -

4.2. COMPUTED TOMOGRAPHY
The SRI SC applied the SRI v3 methodology to Computed Tomography, studying the potential for improvements of the technology, defining a measurement methodology for energy consumption and defining an eco-design goal to develop initiatives with the aim to educate users on how to operate the CT scanners in an environmental friendly way.

4.2.1. INFORMATION TO USERS – ENERGY INFORMATION
The CT energy measurement procedure provides for manufacturers to report the energy consumption of CT systems in the following scenarios:

**Scenario-Off:** This value represents the daily energy consumption when the CT scanner is switched off during the 12h night time (no energy consumption).

**Scenario-LowPower:** This value represents the daily energy consumption when the CT scanner is switched to LowPower during the 12h night time.

**Scenario-Idle:** This value represents the daily energy consumption of the CT when it is left in idle mode for 12h during night time.

Those three values could prove very useful to the user as they underline the difference in energy consumption that can be achieved according to the way the CT scanner is used. They also provide to the user or the purchaser a clear indication on the running cost and how much energy and money can be saved by a correct utilization of the CT.

The following data format has been developed and is recommended for use in product documentation when presenting energy consumption data measured according to the COCIR SRI CT Measurement Methodology. Data should be expressed to the nearest 1 kWh.
Typical Energy Consumption
The typical energy consumption values have been measured according to the COCIR Self-Regulatory Initiative CT Measurement of Energy Consumption, version 1.0

<table>
<thead>
<tr>
<th>Model:</th>
<th>Energy per Day (exemplary values)</th>
<th>Units</th>
<th>Deviation, Justification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scenario-Idle</td>
<td>72</td>
<td>kWh</td>
<td></td>
</tr>
<tr>
<td>Scenario-LowPower</td>
<td>50</td>
<td>kWh</td>
<td></td>
</tr>
<tr>
<td>Scenario-Off</td>
<td>37</td>
<td>kWh</td>
<td></td>
</tr>
</tbody>
</table>

* The system use scenario varies according to customer needs during overnight hours. According to the standard, the system is in active use for 12h during the day and inactive for 12h overnight. The 12h overnight may be in Idle, LowPower, or Off modes with corresponding daily energy consumption variations. Measured values in this table are to be used for economic estimation purposes only. These values do not imply, and are not to be used for, conformance to any clinical or safety requirements.

Additional scenarios can be added by Companies for specific power modes and specific functionalities as long as such scenarios are measured according the COCIR CT Measurement Methodology for LowPower mode.

4.3. CONCLUSION OF THE ULTRASOUND PILOT PROJECT

The Ultrasound pilot project concluded in 2012 as scheduled but the monitoring continued in 2013 as well. As shown in the figure, in 2012 the annual average energy consumption per unit for ultrasound equipment reached 743 kWh/unit against a set target of 691 kWh/unit. A reduction of around 20% in the average annual consumption per model had been achieved compared to the 2005 baseline. In 2013 the average energy consumption per unit decreased significantly to 643 kWh/unit going beyond the set target.

![Average Annual Energy Consumption Per Unit](image)

*Figure: Ultrasound annual energy consumption per unit: market fleet average*

More information on the pilot project are reported in Appendix II.
4.3.1. INTERPRETATION OF DATA (2005/2013)

The analysis of the data shows and confirms the following conclusions:

1. At least two categories of U/S equipment could be identified with very different behaviour. Low energy using U/S (handheld and laptop based) and high energy using U/S (large self-standing hospital equipment). See figure 1 and figure 2.

2. Each category displays a decreasing trend of the annual average energy consumption per unit.

4.3.2. LOW AND HIGH ENERGY USING U/S

Collected data confirms 2 categories with different behaviour can be defined for U/S equipment. This limit of the ultrasound project was already understood and reported previously and integrated in the SRIv2 and in the MRI project (where 3 categories have been identified).

1. Low energy using U/S: handheld and laptop based ultrasound equipment.


As figure 1 below shows, the two categories are subject to different dynamics in sales. High energy U/S sales decreased constantly between 2006 and 2010 when they started to increase again. During the same period, the sales of low energy using U/S show an erratic behaviour.

In particular the increase of High U/S in 2012 together with the strong decrease in sales of Low U/S explains the increase in the average annual energy consumption per unit registered in 2009 and 2012. In 2011 sales of Low U/S increased less than the sales of High U/S therefore increasing the average energy per unit.

![SALES OF LOW ENERGY AND HIGH ENERGY USING U/S](image)

*Figure 1: Sales of high energy using and low energy using ultrasound equipment compared to 2005 baseline*
4.3.3. REDUCTION IN ENERGY CONSUMPTION DUE TO TECHNICAL IMPROVEMENTS

In the COCIR SRI both technical improvements and sales mix account for target achievement. While technical improvements can be controlled by companies, sales can be influenced but not controlled. As shown so far, market demand is hardly predictable and inconstant therefore it could play a critical role in target achievement (positive or negative).

Nonetheless the subdivision of U/S in two different categories allows analysing the energy behaviour of each category.

As clearly shown in figure 2 the average energy consumption of both categories has been following a decreasing trend since 2005 (with some exception). In 2012, High U/S being a mature technology registered a reduction of \(\sqrt{10.5}\%\) while Low U/S, a more recent technology, registered a reduction of \(\sqrt{14.4}\%\).

Figure 2: Fleet average for high and low energy consuming ultrasound products
2013 -

4.4. X-RAY
The SRI SC studied the x-ray modality in 2013. Unlike other imaging equipment in the scope of the SRI there is a big number of companies manufacturing x-ray devices. The actual SRI participating companies cover around 92% of the market for angiography systems, but less than 52% for all the remaining categories.

X-ray devices are quite simple compared to the modalities analyzed so far by the SRI (MRI and CT) and the energy consumption is lower (See appendix V).

Considering the great variety of x-ray categories and different uses (from mobile c-arc to stationary interventional systems), it is impossible to identify a general user behaviour. Nonetheless considering the findings of previous projects it can be expected that most users do not profit of the low power or switch off functions of x-ray devices. This is supported by the study of the Danish Energy Saving Trust “Energy Efficiency in Hospitals and Laboratories”, where the energy consumption of x-ray and user behavior have been investigated.

Already existing Off and LowPower modes, if used properly by users could ensure an energy saving between 50,5% and 64,3% of daily energy consumption (see Appendix V). Nonetheless such options are not widely used by users. The use of Off mode during the night hours and weekend can save on average up to 3,45 MWh per year per equipment.

The COCIR SRI has set an industry goal to develop initiatives with the aim to educate users on how to operate the x-ray scanners in an environmental friendly way.

4.4.1. COCIR “GUIDELINES FOR X-RAY USERS ON SAVING ENERGY”
The development of information material to be disseminated to users can play a significant role in changing user perception and therefore in influencing behaviours. Given the importance of potential savings in the use of x-ray equipment, the SRI SC Committee started the development of a new brochure for x-ray in the course of 2014, which has been disseminated to users and purchasers.

COCIR draft brochure: “COCIR x-ray Guidelines for Users on Saving Energy - Good Environmental Practice”
4.4.2. COCIR “GUIDELINES FOR CT USERS ON SAVING ENERGY”

The development of information material to be disseminated to users can play a significant role in changing user perception and therefore in influencing behaviors. In 2013 COCIR developed the “COCIR Guidelines on energy saving on Computed Tomography – Contribution to healthcare environmental sustainability” which have been officially released in January 2014.

The objective of the Guidelines is not only to provide useful recommendations but also to provide figures on the possible savings achievable with a proper use. Such savings can be quantified for the first time thanks to the COCIR SRI methodology for energy measurement.

The brochure has been disseminated to users and patient organizations:
- European Society of Radiology (ESR): http://www.myesr.org
- European Federation of Nurses Associations (EFN): http://www.efn.be/
- Healthcare Without Harm (HCWH): http://www.noharm.org/
- Head of European Radiological protection Competent Authorities (HERCA): http://www.herca.org

The guidelines have been also distributed in paper version at the European Congress of Radiology (ECR), held in Vienna beginning of March 2014.

2014 -

4.5. REVISION OF THE ULTRASOUND METHODOLOGY

In 2009 the SRI Steering Committee set a target for the Ultrasound Pilot project to achieve by 2012 a 25% reduction in energy consumption compared to 2005, the year chosen as reference. The Pilot concluded in 2012 while the monitoring continued until 2014. The achievements of the pilot are reported in chapter 4.3.

In 2014 the SRI Steering Committee decided to focus again the attention on Ultrasound without waiting for 2016, as scheduled. The pilot methodology for the measurement of the energy consumption of ultrasound equipment was developed in 2009 and was not intended to produce data for comparing different product performances. With the incoming revision of the EU GPP criteria for medical devices, COCIR would like to see a better methodology included in the GPP criteria for U/S equipment.

Therefore, in 2015, the SRI SC released the new “U/S measurement of energy consumption:
methodology” paper which can be freely downloaded from the COCIR website. At the same time the monitoring of the Ultrasound energy consumption has to be halted. In fact, the new methodology is going to change the way U/S devices are measured and therefore the new figures will not be comparable with the old measurements.

The SRI SC discussed in 2015/2016 the feasibility of restarting the monitoring, in particular considering the time needed to companies to re-measure their equipment with the new methodology. In particular for old models this could be expensive and time consuming. The activity may be limited to newly released models. New environmental aspects such as use of resources may be evaluated as well considering the continuous reduction in size of U/S devices.

4.6. COCIR “GUIDELINES FOR MRI USERS ON SAVING ENERGY”

Following the findings of the CT project, COCIR realized that the user behaviour has an important influence on the energy consumption of MRI.

Scenarios have been defined:
- **Scenario-Off**: the MRI scanner is turned in Off mode during the 12h night time.
- **Scenario ready-to-scan**: the MRI is always in ready-to-scan mode.

Considering that during the weekend the scanner can be in Off-mode or Ready-to-scan mode for 24 hours, the use of the Off-mode allows for saving up to 21,8% of the annual energy consumption which on the average corresponds to 29,2 MWh/y.

The development of information material to be disseminated to users can play a significant role in changing user perception and therefore in influencing behaviors. Given the importance of potential savings, the SRI SC Committee developed a new brochure for MRI in the course of 2014 which has been disseminated to users and purchasers.

---

4 The MRI cryo-cooler cannot be stopped therefore there Off-mode has a significant energy consumption
4.7. NUCLEAR MEDICAL IMAGING
The COCIR SRI focuses on PET and SPECT nuclear imaging modalities. Combined technologies such as PET-CT, SPECT-CT, PET-MRI are not considered for the complexity of the technology, and the recent introduction on the market (2010 for the first installed PET-MRI).

The initial analysis of medical nuclear images shows that PET is not sold anymore as single technology but always in combination with CT or MRI, due to huge increment in clinical value of adding precision of anatomic localization to functional imaging.

SPECT is sold in small numbers (116 units in 2013) and presents very low energy consumption, comparable to X-ray systems. The comparison of the average energy consumption of SPECT (15 kWh/d) with other modalities such as MRI (250/300 kWh/d or CT 70kWh/d) makes clear the limited relevance of SPECT energy consumption.

For the above mentioned reasons, and considering that nuclear imaging is not included in the EU GPP criteria, the SRI SC decided not to proceed with the development of a measurement methodology for SPECT or PET but to give priority in 2014/2015 to the definition of the U/S measurement methodology, giving also the impending revision of the EU GPP criteria for EEE in the medical sector.

Considering the power usage stability of PET and SPECT in different modes, and the limited potential savings achievable by users the SRI SC did not deem necessary to develop a COCIR Guideline to energy saving for nuclear medicine at least for this year.
4.8. MAGNETIC RESONANCE ACHIEVEMENTS

The SRI Steering Committee calculated the average annual energy consumption for new products put on the market for the year 2016. The data is used to assess achievements compared to the baseline 2011 (see Appendix III for additional details on methodology and calculations).

As shown by table 3, in 2016 the daily average energy consumption per unit for MRI equipment decreased to 173,25 kWh/unit showing a 23% reduction compared to 2011 and a 2% compared to 2015.

<table>
<thead>
<tr>
<th>Sold units</th>
<th>Total daily energy consumption (kWh)</th>
<th>Average daily energy consumption per unit (kWh/d)</th>
<th>Beyond BAU (kWh/d)</th>
<th>BAU (kWh/d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2011</td>
<td>✓ 394</td>
<td>✓ 89.011</td>
<td>✓ 225,92</td>
<td>✓ 225,61</td>
</tr>
<tr>
<td>2012</td>
<td>✓ 446</td>
<td>✓ 100.038</td>
<td>✓ 224,30</td>
<td>✓ 225,30</td>
</tr>
<tr>
<td>2013</td>
<td>✓ 454</td>
<td>✓ 95.148</td>
<td>✓ 209,58</td>
<td>✓ 225,00</td>
</tr>
<tr>
<td>2014</td>
<td>✓ 513</td>
<td>✓ 95.572</td>
<td>✓ 186,30</td>
<td>✓ 224,69</td>
</tr>
<tr>
<td>2015</td>
<td>✓ 604</td>
<td>✓ 106.855</td>
<td>✓ 176,91</td>
<td>✓ 224,38</td>
</tr>
<tr>
<td>2016</td>
<td>✓ 680</td>
<td>✓ 117.808</td>
<td>✓ 173,25</td>
<td>✓ 224,07</td>
</tr>
<tr>
<td>2017</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4.8.1. TOTAL ENERGY SAVINGS

When the MRI project was launched, it was estimated that SRI could save, by 2017, compared with the BAU baseline, around 7,459 kWh per unit sold according to the Beyond BAU scenario, equivalent to more than 2,3 tons of CO₂ per year per unit.

In 2016 it is possible to calculate the cumulative effective savings of energy, comparing the achievements with the baseline scenario.

<table>
<thead>
<tr>
<th>UNITS</th>
<th>YEAR</th>
<th>BAU kWh/d</th>
<th>kWh/u</th>
<th>SRI kWh/d</th>
<th>kWh/u</th>
<th>MWh/y</th>
</tr>
</thead>
<tbody>
<tr>
<td>394,0</td>
<td>2011</td>
<td>89.010</td>
<td>225,75</td>
<td>89.010</td>
<td>225,91</td>
<td>0</td>
</tr>
<tr>
<td>446,0</td>
<td>2012</td>
<td>100.684</td>
<td>225,75</td>
<td>100.036</td>
<td>224,30</td>
<td>842</td>
</tr>
<tr>
<td>454,0</td>
<td>2013</td>
<td>102.490</td>
<td>225,75</td>
<td>95.147</td>
<td>209,57</td>
<td>7.637</td>
</tr>
<tr>
<td>513,0</td>
<td>2014</td>
<td>115.810</td>
<td>225,75</td>
<td>95.126</td>
<td>218,43</td>
<td>16.133</td>
</tr>
<tr>
<td>604,0</td>
<td>2015</td>
<td>136.353</td>
<td>225,75</td>
<td>106.855</td>
<td>176,91</td>
<td>15.338</td>
</tr>
<tr>
<td>680,0</td>
<td>2016</td>
<td>153.510</td>
<td>225,75</td>
<td>117.808</td>
<td>173,25</td>
<td>9.282</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>49.235</td>
</tr>
</tbody>
</table>

7 Values are slightly different from the ones in SRI Report 2012 due to a retrospective re-categorization of products by one company causing a variation of SRI sales and related energy consumption data of 0.15% in 2011 and 0.13% in 2012 compared to the values reported in previous SRI Status Reports.

8 Sold units data provided by companies for each model placed on the market in the calendar year.

9 Measured energy consumption data provided by companies for each model placed on the market in the calendar year.

10 Assuming 5 days per week, 52 weeks per year.

The SRI allowed between 2010 and 2016 the saving of 49 GWh, equivalent to 15.2 tons of CO2 (a small town of 12,000 households).

4.8.2. INTERPRETATION OF THE DATA

The fleet average energy consumption continued the reduction trend in 2016. While fluctuations in the fleet average may be due to a different sales mix, the drops in energy use are mostly due to the increased performance of participating companies, as shown by figure 2.1.

Figure 2.1: trend in companies’ fleet average daily energy consumption
4.9. CONCEPT METHODOLOGY TO MEASURE CIRCULAR ECONOMY

In 2016, at the 5th Annual Forum on the COCIR SRI, COCIR presented its conclusions about reuse and refurbishment activities being the most significant contribution to reducing environmental impact of medical imaging devices. In 2015/2016 COCIR worked on a first methodology to estimate the circularity of the economic model.

The European Union is aiming at moving the current “linear” model towards a circular one by means of modification of current legislation and introduction of new one. But moving from a linear model to a circular one is not a linear process. While it is clear that a perfect circular economy cannot be achieved, the degree of success depends on the implementation of all possible measures to prevent waste generation and use of resources from the design to the end of life phase, and to boost recycling of waste. The system complexity therefore increases linearly with the number of feedback loops which bring resources back to use diverting them from landfill, which is considered the end of the cycle.

From a linear economy model

To a circular model

The methodology drafted in Appendix VII aims at defining an index which can be later used for monitoring improvements. Additional work is needed to shape the methodology in such a way that the calculation of the index is based on data which can be obtained by companies with good approximation.
2017 -

4.10. FEASIBILITY EVALUATION
In 2016 the SRI SC launched a project to evaluated the feasibility of determining or estimating the data required to calculate the Circularity Index defined in 2016 (see Appendix VII). A questionnaire was sent to all SRI companies in the refurbishment business.

4.10.1. OBJECTIVE OF THE QUESTIONNAIRE
The objective was to collect expert opinions on the feasibility of collecting the required data, and suggestions on estimations/simplifications. Companies were required to answer:
- Y – The data is available or is readily available
- N – The data is not available and it is impossible to estimate it
- Can be estimated – the data is not available but can be estimated

4.10.2. REQUIRED DATA
The methodology aim is to calculate an index (C.I. Circularity Index) characterizing the "circularity" of the model

\[
C.I. = \frac{RM + PM + RE + RP}{POM} \times 100
\]

Where:
- RM: Mass of recycled materials
- PM: Mass of parts re-used in manufacturing of new equipment
- RE: Mass of refurbished equipment
- RP: Mass of reused parts in repair and maintenance
- POM: Mass of new products and parts placed on the market
4.10.3. CONSOLIDATED ANSWERS

**RM: Mass of recycled materials**

|                        | Feasibility EU | Feasibility Global | Comments                                                       |
|------------------------|----------------|--------------------|                                                               |
| Equipment sent to recycling per year | Y              | Y                  | Available from the reporting according to the WEEE Directive  |
| Waste from refurbishment and other activities per year | Y              | Y                  | Available from waste management systems                       |
| Recycling rate         | C              | C                  | Available from recyclers                                      |

**RP: Mass of reused parts in repair and maintenance**

<table>
<thead>
<tr>
<th></th>
<th>Feasibility EU</th>
<th>Feasibility Global</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Mass of recovered/refurbished (R/R) spare parts per year</td>
<td>C</td>
<td>C</td>
<td>The split between EU and Global might be very difficult and might only be a “good guess”</td>
</tr>
<tr>
<td>Mass of (R/R) parts used for refurbishment of used MDs per year</td>
<td>C</td>
<td>C</td>
<td></td>
</tr>
<tr>
<td>Mass of (R/R) parts used in other activities (or mass of parts sent to waste in repair and maintenance activities) per year</td>
<td>C</td>
<td>C</td>
<td>Isn’t that the difference between the total mass and the mass used for Refurbishment?</td>
</tr>
</tbody>
</table>

*Note: While it may be possible to estimate the number of parts used in repair and maintenance of the installed base it is more difficult to know the mass as the weight of parts is not necessarily recorded and due to the heterogeneity, an average weight is meaningless. For parts discarded as waste, the weight should be known. The complexity is represented by the numerous service providers who normally take care of installed equipment and which, sometimes are not related to the OEM.*

**Total mass of reused parts in production of new equipment**

Since July 2014 and with the expiration of exemption 31 on 6 November 2017 (substituted by exemption 31a), this activity will be mostly forbidden and therefore close to zero (with the exception of RoHS compliant spare parts).

**RE: Total mass of refurbished equipment placed on the market**

<table>
<thead>
<tr>
<th></th>
<th>Feasibility EU</th>
<th>Feasibility Global</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Units of recovered/refurbished equipment placed on the market</td>
<td>Y</td>
<td>Y</td>
<td>Equipment “as is” can be difficult to estimate</td>
</tr>
<tr>
<td>Average Weight</td>
<td>C</td>
<td>C</td>
<td></td>
</tr>
</tbody>
</table>
POM: Total Mass of new equipment and new parts placed on the market

<table>
<thead>
<tr>
<th></th>
<th>Feasibility EU</th>
<th>Feasibility Global</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Mass of new equipment placed on the market per year</td>
<td>Y</td>
<td>Y</td>
<td>From sales record, can be estimated based on sales and average product weight</td>
</tr>
<tr>
<td>Total mass of new parts used for refurbishment</td>
<td>C/N</td>
<td>C/N</td>
<td>The split between EU and Global can only be guessed. In addition it is very time-consuming to get the data.</td>
</tr>
<tr>
<td>Total mass of new parts used for repair or other activities</td>
<td>C/N</td>
<td>C/N</td>
<td>The split between EU and Global can only be guessed. We could make some estimates, but may take some time to build up the proper way of working</td>
</tr>
</tbody>
</table>

4.11. SUB-INDEXES

The results of the feasibility study prove that certain data required to calculate the index would require additional work to be estimated. For this reason, the SRI SC decided to launch in 2016/2017 a data collection for the definition of a sub-index.

\[ C.I. = \frac{RE + PM + RM + RP}{POME + POMp} * 100 \]

4.11.1. RE/POME

Tons of refurbished equipment sold in 2016 / Tons of new equipment sold in 2016. This sub-index measures the ratio between the tons of refurbished equipment and the tons of new equipment placed on the market during the year.

Assuming the average weight of new equipment and refurbished ones to be the same, the RE/POME ratio can be calculated using units as the average weights delete each other.

<table>
<thead>
<tr>
<th>2016</th>
<th>UNITS SOLD</th>
<th>UNITS REFURB</th>
<th>RE/POME %</th>
</tr>
</thead>
<tbody>
<tr>
<td>MRI</td>
<td>871</td>
<td>46</td>
<td>5,3%</td>
</tr>
<tr>
<td>CT</td>
<td>993</td>
<td>52</td>
<td>5,2%</td>
</tr>
<tr>
<td>X-RAY</td>
<td>2942</td>
<td>152</td>
<td>5,1%</td>
</tr>
<tr>
<td>TOTAL</td>
<td></td>
<td></td>
<td>5,2%</td>
</tr>
</tbody>
</table>

Data from the COCIR SHARE market statistics tool

Considering that the values for the 3 modalities are virtually the same, it is possible to assume the total ratio even if the average weight for CT, MRI and X-RAY is different.
4.11.2. REUSE OF PARTS

The data collection showed that COCIR companies collect and repair/refurbish around 6400 metric tons of parts per year globally. The weight of refurbished equipment (MRI+CT+X-RAY+Ultrasound) accounts for less than 1000 tons per year in EU (estimated 2900 globally). This shows that reuse and refurbishment of parts is one of the main elements of circular economy in the medical imaging devices sector.

![Pie chart showing reuse of medical imaging devices in 2016](image)

Data source: internal data compilation based on a survey of 4 companies involved in refurbishment activities
APPENDIX I

1. SRI GENERIC METHODOLOGY
In 2010 the SRI Steering Committee started to apply the methodology first two steps to all the modalities in scope to identify a priority list.

1.1. GATHER BASELINE DATA FOR ALL MODALITIES IN SCOPE (STEP 1)
Baseline data was gathered according to a specific template (see SRI Methodology V3 Appendixes). According to the product portfolio of the SRI members (see table 1) data has been delivered from the following companies:
- GE Healthcare
- Philips Healthcare
- Siemens Healthcare
- Toshiba Medical Systems Corporation
For the following modalities and sub-modalities:
- Computed tomography (CT)
- Magnetic Resonance Imaging (MRI)
- Nuclear Medicine
  - NM Conventional
  - NM PET
- X-Ray
  - X-Ray Angio
  - X-Ray Fluoro
  - X-Ray Radio
  - X-Ray Mammography
  - X-Ray Surgery

1.2. PRIORITIZATION AND SELECTION OF THE NEXT MODALITY (STEP 2)
The company LCA data from STEP 1 have been consolidated by the SC Secretariat after a plausibility check.

The SC Secretariat calculated two rankings with the provided LCA data:
- The first ranking weights the environmental loads (delivered by the companies in STEP 1) with current units sold in EU (based on SHARE data).
- The second ranking is based on the expert judgements on factors such as feasible technological developments and the complexity of the product, as well as on sales forecasts of the next generation equipment.

The final priority list, see table 5, that is used to select the next modalities is obtained averaging the two previously calculated rankings.

12 Only already available LCA data has been provided. According to SRIv2 methodology Step 1 and 2 do not require LCA data to be provided by all the participating companies.
13 SHARE is COCIR’s internal market statistics data base.
Table 5: Priority list

<table>
<thead>
<tr>
<th>Modality</th>
<th>Environmental loads ranking 2009</th>
<th>Risk Assessment ranking 20xx*</th>
<th>Average Ranking</th>
<th>Final Ranking</th>
</tr>
</thead>
<tbody>
<tr>
<td>MRI</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>CT</td>
<td>3</td>
<td>2</td>
<td>2,5</td>
<td>2</td>
</tr>
<tr>
<td>X-Ray</td>
<td>2</td>
<td>3</td>
<td>2,5</td>
<td>3</td>
</tr>
<tr>
<td>Nuclear Medicine</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

*depends on the typical modality innovation cycle

CT has been chosen as second one as the European market coverage of the participating companies is around 100%, while for X-ray it is about 80%. The higher the market coverage, the higher the reduction of environmental impacts that the SRI Initiative could achieve.

According to table 6, the SRI Initiative will focus on the listed modalities in the given sequence, at least one each year. Thus, MRI was targeted as the first modality under the SRI methodology and CT as second one.

Table 6: Timetable for targeting new modalities

<table>
<thead>
<tr>
<th></th>
<th>2011</th>
<th>2012</th>
<th>2013</th>
<th>2014</th>
<th>2015</th>
</tr>
</thead>
<tbody>
<tr>
<td>MRI</td>
<td>♤</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CT</td>
<td></td>
<td>♤</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>X-Ray</td>
<td></td>
<td>♤</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nuclear Medicine</td>
<td></td>
<td></td>
<td>♤</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ultrasound</td>
<td></td>
<td></td>
<td></td>
<td>♤</td>
<td></td>
</tr>
</tbody>
</table>

1.3. IDENTIFICATION OF SIGNIFICANT ENVIRONMENTAL ASPECT (STEP 3)

Every year a new part is added with a new modality, following the order of the priority list. Each new part for the specific modality includes data on the identification of the top environmental aspect.
APPENDIX II

2. ULTRASOUND IMAGING EQUIPMENT

2.1. MEASUREMENT OF THE ENERGY CONSUMPTION: 2015

In 2014 the SRI Steering Committee decided to focus again the attention on Ultrasound imaging equipment instead of waiting for 2016, as scheduled. The pilot methodology for the measurement of the energy consumption of ultrasound equipment was developed in 2009 and was not intended to produce data for comparing different product performances. COCIR developed a better methodology in particular for battery powered equipment.

2.1.1. SCOPE

The methodology is designed to work for all ultrasound equipment, given that the energy consumption in ready-to-scan mode and scan mode are virtually the same. Certain high-end models are able to perform real-time 3D imaging for some specific examinations (i.e. obstetric). In those particular examinations the power usage during scan can be 20% higher than the power in ready-to-scan modes. Considering the limited numbers of such equipment and of examinations, such models are not covered by this methodology. To measure the energy usage of such models, scan conditions and parameters should be defined to allow the measurement of the energy consumption.

2.1.2. THE METHODOLOGY IN BRIEF

The following functioning modes have been defined for ultrasound equipment:

<table>
<thead>
<tr>
<th>Mode</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off:</td>
<td>The system is shut down according to the user manual, plugged into mains, AC mains ON</td>
</tr>
<tr>
<td>Stand-by</td>
<td>The system is configured in the lowest possible energy consuming state, according to the user manual</td>
</tr>
<tr>
<td>Ready-to-scan</td>
<td>The system is fully powered and ready to acquire image</td>
</tr>
<tr>
<td>Scan</td>
<td>The system is acquiring image. This mode includes ultrasound generation and real-time image rendering</td>
</tr>
</tbody>
</table>

The power usage is measured in the different modes and the daily energy usage calculated according to 3 different scenarios.

<table>
<thead>
<tr>
<th>Scenarios and distribution of time in functioning modes</th>
<th>Time in Off mode (h)</th>
<th>Time in Stand-by mode (h)</th>
<th>Time in Ready-to-scan mode (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scenario Off</td>
<td>12</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Scenario Stand-by</td>
<td>0</td>
<td>18</td>
<td>6</td>
</tr>
<tr>
<td>Scenario Ready-to-scan</td>
<td>0</td>
<td>0</td>
<td>24</td>
</tr>
</tbody>
</table>

The energy consumption expressed in different scenarios is useful as it allows the immediate comparison of the energy consumption related to different ways of using the equipment.

The monitoring of the Ultrasound energy consumption has to be halted concluding the Pilot Project. In fact the new methodology is going to change the way U/S devices are measured and therefore the new figures will not be comparable with the old measurements.
2.2. ULTRASOUND IMAGING EQUIPMENT PILOT PROJECT (2009/2012): CONCLUDED

When COCIR and the participating companies started to develop the Self-Regulatory Initiative, the methodology and approach was a new terrain for all. Participating companies initiated a first non-standardised review of the products in scope to establish the product of initial focus for a pilot, to develop the industry baseline for energy trending, and to establish targets and timing. The final choice of ultrasound imaging equipment was based on the following reasons:

- Ultrasound equipment is manufactured by the majority of the COCIR participating companies in the Self-Regulatory Initiative: Hitachi Medical Systems Europe (ex-Aloka)\(^\text{14}\), GE Healthcare, Hitachi Medical Systems Europe, Samsung (ex-Medison)\(^\text{15}\), Philips Healthcare, Siemens Healthcare and Toshiba medical Systems Europe. The inclusion of many manufacturers could steep the learning curve.
- COCIR members have a good understanding of environmental aspects and opportunities to reduce environmental impacts for ultrasound equipment. Most of the manufacturers are engaged in Life Cycle Analysis projects and focus on the ecodesign of their products.
- Ultrasound equipment is much less complex, including fewer components compared to other modalities in the medical imaging sector. Taking ultrasound as example was easier and faster to learn and to develop a methodology, to establish company internal processes to assess environmental aspects, create targets, and to change technologies.

2.3. GENERAL DESCRIPTION OF ULTRASOUND EQUIPMENT

Ultrasound is an imaging technique used to visualize subcutaneous body structures including tendons, muscles, joints, vessels and internal organs for possible pathology or lesions. Obstetric ultrasound is commonly used during pregnancy to check on the development of the fetus.

Ultrasound uses a piezoelectric transducer encased in a probe to send pulses of sound into the body. The sound wave is partially reflected at each point in the body where a tissue interface results in a change in density. The time it takes for the echo to travel back to the transducer is measured and used to calculate the depth of the tissue interface causing the echo. The greater the difference in density, the larger the echo is.

The sound is focused either by the shape of the transducer, a lens in front of the transducer, or a complex set of control pulses from the ultrasound scanner machine. This focusing produces an arc-shaped sound wave from the face of the transducer. The wave travels into the body and comes into focus at a desired depth.

Typical ultrasound scanners operate in the frequency range of 2 to 18 megahertz, hundreds of times greater than the limit of human hearing. The choice of frequency is a trade-off between spatial resolution of the image and imaging depth. Superficial structures such as muscles, tendons, testes, breasts and the neonatal brain are imaged at a higher frequency (7-18 MHz), which provides better axial and lateral resolution. Deeper structures such as liver and kidney are imaged at a lower frequency 1-6 MHz with lower axial and lateral resolution but greater penetration.

\(^{14}\) Aloka has been acquired by Hitachi in 2012
\(^{15}\) Medison has been acquired by Samsung in 2012
2.4. MARKET COVERAGE

The 7 companies \(^{16}\) participating in the ultrasound pilot project had a total turnover in Europe in 2013 of 737,4\(^{17}\) million euros covering around 82\(^{18}\) \% of the European market.

**Table 4: Ultrasound - EU\(^{19}\) market data**

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Ultrasound (US)</td>
<td>(788, \text{M€})</td>
<td>(740, \text{M€})</td>
<td>(737,4, \text{M€})</td>
<td>82%</td>
</tr>
</tbody>
</table>

2.5. SRIV1 METHODOLOGY FOR ULTRASOUND

Participating companies developed a generic process to be followed for the pilot ultrasound. A detailed description of each process phase is described in the SRIV1.

2.6. TARGET SETTING

The industry SRI SC set a target to reduce in 2012 the average annual energy consumption per unit of new ultrasound products placed on the market by 25\% compared to the 2005 baseline. The target was set on the basis of an expert judgment on realistic feasibility to improve the already existing ecodesign programs.

To reach the 25\% target, according to table 2, participating companies reduced the average annual energy consumption per unit by 14,5\% from 2009 to 2012. The average annual energy consumption per unit has been reduced despite new products had increased functionality and delivered even more healthcare benefits to patients.

The target to reduce average annual energy consumption of new ultrasound products placed on the market by 14,5\% between 2009 and 2012 translates to a reduction in average annual energy consumption from 809 kWh per unit per year in 2009 down to 691 kWh per unit per year in 2012.

Participating companies achieved the target by setting the following objectives:

- Increased focus on ecodesign in the product design and development process. For example, considering the use of the International Standard IEC 60601-1-9: Environmentally Conscious Design of Medical Electrical Equipment
- Specify and design product components and parts with much less energy consumption
- Using new technologies (e.g. Green IT equipment)

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\(^{16}\) SRI Member: GE Healthcare, Hitachi Medical Systems Europe, Philips Healthcare, Siemens Healthcare, Toshiba Medical Systems Europe, Hitachi-Aloka, Samsung.

\(^{17}\) OCIR SHARE internal market statistics data base. Data are based on the fiscal year.

\(^{18}\) Estimation provided by companies based on SHARE data

\(^{19}\) OCIR Imaging Market Statistics source (SHARE). Countries included: Estonia, Latvia, Lithuania, Bosnia, Bulgaria, Croatia, Czech Republic, Hungary, Poland, Romania, Serbia, Slovakia, Slovenia, Ukraine, Portugal, Spain, Denmark, Finland, Norway, Sweden, Ireland, UK, Austria, Belgium, France, Germany, Greece, Italy, Netherlands, Switzerland, Albania, Macedonia.
3. MAGNETIC RESONANCE IMAGING EQUIPMENT

3.1. GENERAL DESCRIPTION OF MAGNETIC RESONANCE EQUIPMENT

Magnetic resonance imaging (MRI) is a medical imaging technique used in radiology to visualize detailed internal structures of the human body. MRI makes use of the property of magnetic resonance of nuclei to create medical diagnostic images.

An MRI machine utilizes superconductor technology by using liquid Helium (below 4.2 Kelvin) to create a powerful magnetic field to align the magnetization of atoms within the body. Radio frequency waves are used to systematically alter the alignment of this magnetization. This causes the nuclei to produce a rotating magnetic field detectable by the scanner. Very powerful magnetic field gradients are needed to cause nuclei at different locations to rotate at different speeds providing the necessary 3-D spatial information. The information collected is manipulated with high speed mathematical formulas to generate extremely detailed medical diagnostic images.

MRI provides excellent contrast between the different soft tissues of the body, which makes it especially useful in imaging the brain, muscles, internal organs, and cancers. Compared with other medical imaging techniques such as computed tomography (CT) or X-rays, MRI uses no ionizing radiation.

MRI technologies

MRI equipment uses two different technologies to generate the required magnetic field strength that could vary from 0.35 Tesla up to 7 Tesla or even more.

Permanent magnet: permanent magnets are used to generate magnetic field up to 1.2 Tesla. Commonly such models are equipped with non-cylindrical magnets allowing more patient comfort. Non cylindrical magnet MRIs are called “Open MRI”.

Superconductive magnet: superconductive electromagnets, cryo-cooled to 4 Kelvin using liquid helium, are used to generate magnetic fields up to 7 Tesla or more. The boiled helium is re-condensed by a cryo-cooler (Gifford-McMahon or pulse tube). The cryo-cooling system cannot be switched off except in case of emergency. This causes the helium to boil off and get lost. Normally superconductive MRIs use cylindrical magnets but sometimes open magnets.

Figure 4: Open MRI and cylindrical MRI
Magnetic field strength
The strength of the magnetic field and the power of the gradient coil and the RF senders determine the quality and resolution of the image. High end machines for hospital use are equipped with 3 Tesla magnets. Higher fields equipment, up to 7 Tesla, are actually under development and test and used only for research purposes.

Bore size
The bore diameter is important for patient comfort. Patient suffering from claustrophobia could experience better comfort in larger bores. Moreover, large bores allow the examination of “big” patients suffering from obesity. Nonetheless larger bore size requires the use of more powerful and energy consuming magnet systems and gradient coils, as the field strength decreases with the distance.

3.2. MODES
Three modes have been defined for MRI equipment.

Off mode:
The MRI is in the lowest user selectable power state. In superconductive MRIs the magnet needs to be cooled permanently. Therefore, the cooling circuitry and the magnet supervision needs to be active.

Ready-to-scan mode:
The MRI is on and ready to acquire an image. All modules are active. However, neither gradient pulses nor radio frequency waves are sent or received. The computing system may calculate and display images from raw data previously acquired.

Scan mode:
The MRI is actively scanning the patient by sending high frequency waves as well as gradient pulses and reading the resulting variations in the magnetic field. The computing system acquires the corresponding data and calculates and displays images.

The power consumption of MRI in the three modes is represented in figure 5.

![Image](image_url)

**Figure 5:** Exemplary power consumption of MRI

---

20 In case the magnet cooling system is switched off, the helium slowly boils and it is released. The released helium is lost and needs to be replaced by liquid helium. This implies the corresponding cooling and transporting efforts.
3.3. POWER CONSUMPTION
The measurements performed on all models allowed to determine the energy consumption of MRI equipment in the different operating modes, according to the defined use scenario (Appendix III.II.IV).
Even if the variability between different MRI is relevant, the following average values can be identified equipment:

<table>
<thead>
<tr>
<th>MODE</th>
<th>Average Power Consumption (kW)</th>
<th>Average distribution of daily energy consumption %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off</td>
<td>9,3</td>
<td>34</td>
</tr>
<tr>
<td>Ready to scan</td>
<td>14,6</td>
<td>34</td>
</tr>
<tr>
<td>Scan</td>
<td>22,3</td>
<td>32</td>
</tr>
</tbody>
</table>

The power consumption in scan mode cannot be easily measured as it is different for each sequence and moreover it varies extremely during the same sequence as shown by figure 6. For each sequence the average power consumption has to be derived.

![Figure 6: Power consumption for different sequences in abdomen examination](image)

**Figure 6**: Power consumption for different sequences in abdomen examination
3.4. MARKET DATA

The 5 companies participating in the SRI for the MRI sector represent a total turnover in Europe of 804 million euros in 2016, covering about 100% of the European market.

Table 7: MRI ~ EU24 market data

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnetic Resonance Imaging (MRI)</td>
<td>€677 M€</td>
<td>€762 M€</td>
<td>€804 M€</td>
<td>100%</td>
</tr>
</tbody>
</table>

3.5. IDENTIFICATION OF THE MOST SIGNIFICANT ENVIRONMENTAL ASPECT FOR MRI

Magnetic resonance Imaging equipment has been chosen by the Steering Committee as the first modality to be targeted on the base of Step 1 and Step 2 of the methodology as shown in table 5.

According to Step 3 of the methodology, data provided by companies are used to rank the different environmental aspects. Table 8 shows that energy consumption during the product lifecycle use phase has been identified as the top environmental aspect.

Table 8: Identification of most significant environmental aspects

<table>
<thead>
<tr>
<th>Identification of most significant environmental aspect</th>
<th>Aspects</th>
<th>Average internal ranking</th>
<th>Final COCIR Ranking</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Energy use</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Non ferrous metals</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Ferrous alloys</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Helium consumption</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Magnet metals</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>Copper in Gradient coil</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Copper: end of life</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>Copper: production</td>
<td>3</td>
<td>7</td>
</tr>
</tbody>
</table>

21 GE Healthcare, Hitachi Medical Systems Europe, Philips Healthcare, Siemens Healthcare, Toshiba Medical Systems Europe
22 COCIR SHARE internal market statistics data base
23 Estimation provided by COCIR companies based on 2016 SHARE data
24 COCIR Imaging Market Statistics source (SHARE). Countries included: Estonia, Latvia, Lithuania, Bosnia, Bulgaria, Croatia, Czech Republic, Hungary, Poland, Romania, Serbia, Slovakia, Slovenia, Ukraine, Portugal, Spain, Denmark, Finland, Norway, Sweden, Ireland, UK, Austria, Belgium, France, Germany, Greece, Italy, Netherlands, Switzerland, Albania, Macedonia.
3.6. COMPLEMENTARY DOCUMENTATION

This report is completed by the following documentation:

1. Magnetic resonance Equipment (MRI) - Study on the potential for environmental improvement by the aspect of energy efficiency

The SRI Steering Committee hired in July 2011 an external consultant with long experience in the field of ecodesign, PE International, to study the potential for improvement of MRI equipment with regard to energy efficiency. The study analyses MRI energy consumption, the allocation of power usage in the different modules during off, ready-to-scan and scan mode and technological solutions to improve the efficiency.

Results of the study are used as input for Step 4 of the SRI methodology for setting the ecodesign target for MRI.

2. Magnetic resonance Equipment (MRI) - Measurement of energy consumption

The SRI Steering Committee mandated in October 2010 an Expert Working Group on MRI with the objective to develop a methodology to measure the energy consumption as there are no recognized standards at the moment. The measurement methodology allows company to measure the energy consumption of MRI on a common basis providing comparable data that are used in Step 4 of the SRI methodology.

3.7. MEASUREMENT OF THE ENERGY CONSUMPTION

The SRI Steering Committee started in October 2010 to develop a methodology to measure the energy consumption of MRI equipment, as there are no available standards that could be used.

The methodology ensures that:
- Results of the measurements are comparable and repeatable
- The measured energy consumption has a strict relation with the real energy consumption in everyday hospital use. Such information is important for hospitals and clinics to understand the real running cost of MRI to plan and allocate correctly their budget
- The procedure does not involve disproportionate costs or resources
- The measurement results allow the determination of all the information that could be considered relevant for the SRI target setting process but could also be useful for MRI users.

A first methodology was defined in May 2011 and participating companies started a measurement campaign providing a first set of 5 measured machines. After a deep analysis of the data the methodology was simplified by introducing average values for the ready-to-scan mode durations (see the "Magnetic Resonance – Measurement of energy consumption" document for additional information). Participating companies measured all their MRI models according to the new methodology.

The study on the MRI potential for improvement showed that the energy use in scan mode could not be reduced due to physics of the process which requires a certain amount of energy (see "Magnetic resonance Equipment (MRI) - Study on the potential for environmental improvement by the aspect of energy efficiency").

For the above mentioned reason, the SRI adopted a simplified version of the methodology without measuring the energy consumption in scan modem (for additional information see Appendix III.V).

The methodology has been finalized in February 2012 and is available for download at COCIR website.
3.7.1. MEASURING THE ENERGY CONSUMPTION

The energy consumption could normally be calculated by summing the energy consumption in each mode, calculated multiplying the power consumption for each mode for the relative duration:

\[
\text{Energy use} = T_{\text{off}} \times P_{\text{off}} + T_{\text{ready-to-scan}} \times P_{\text{ready-to-scan}} + T_{\text{scan}} \times P_{\text{scan}}
\]

The average power draw in off mode, servicing mode and ready-to-scan mode can be easily measured. For MRI the following elements are unknown:

- \( T_{\text{ready-to-scan}} \): Duration of ready to scan mode
- \( T_{\text{scan}} \): Duration of scan mode
- \( P_{\text{scan}} \): Power consumption in scan mode

Those durations depend very much on which examination is performed, the scan speed of the machine and the administrative operations to be performed by doctors during the examination (patient preparation, data input, data archiving, patient positioning, etc.). While for off mode it is easy to set an average value according to hospital practices, setting average values for the remaining two modes would not allow to take into consideration a very important factor, the “productivity” of the MRI, the number of patients that can be examined per day (see Appendix II.V).

3.7.2. SYSTEM BOUNDARIES

The SRI SC defined the system boundaries (which modules should be included in the measurement and which not) for the measurement of MRI equipment.

**In:** All system-critical items needed to perform a basic scan, e.g. gradient amplifiers, RF unit, MR coils needed for the specific measurements, reconstruction engine(s), required electronics such power supplies, controllers, console/computer, cryogen compressor, water heat exchanger, patient table, magnet, helium-conservation equipment.

**Out:** Any equipment and accessories beyond basic product offering and not required for a basic scan, or customer-provided equipment, e.g. optional MR coils, patient vital signs accessories, facility-provided cooling water equipment and hardware for advanced medical applications.

3.7.3. EQUIPMENT CONFIGURATION

To allow comparability of the measurements the SRI SC identified ranges for the values of the most relevant parameters for each one of the defined sequences having an impact on the energy consumption:

- Number of slices
- Field of view
- Slice thickness
- Resolution
- Bandwidth
- Sequence duration

As shown in table 9, a set of parameter has been defined for each sequence. The values have been determined on the basis of the experience of companies’ experts as the most commonly used in hospital practice.

Moreover, the values have been validated according to the following documentation:

- the German “Guidelines of the Federal Medical Council for Quality Assurance of magnetic resonance imaging” (BÄK)
- and the “guidelines on criteria for quality assessment in nuclear magnetic resonance imaging pursuant to § 136 SGB V i.V.m. § 92 SGB V, Section 1 of the Federal Committee of Physicians and Sickness Funds (Quality assessment guidelines for magnetic resonance imaging)
For each parameter and for each sequence the minimum or maximum value is indicated in the table.

Table 9: Abstract from the configuration parameters table. The complete table is available in the "MRI – Measurement of energy consumption" document.

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Number of Slices</th>
<th>FOV (mm²)</th>
<th>Slo Thk (mm)</th>
<th>Resolution</th>
<th>Bandwidth (Hz/Px Range)</th>
<th>Seq Req</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Minimum</td>
<td>Max</td>
<td>Min</td>
<td>Max</td>
<td>Min</td>
<td>Max</td>
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<td>200</td>
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<td>290</td>
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<td>0,7</td>
<td>151,0</td>
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<td>500 x 225</td>
<td>5</td>
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<td>222,0</td>
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<td></td>
<td>ep2d0_diff_scan_trace_p2</td>
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<td>1,3</td>
<td>400</td>
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<td>1,4</td>
<td>600</td>
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<td>3</td>
<td>450</td>
<td>8</td>
<td>1,8</td>
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<td></td>
<td>l1_loc.tr.023</td>
<td>3</td>
<td>300</td>
<td>5,0</td>
<td>1,0</td>
<td>250</td>
</tr>
</tbody>
</table>

3.7.4. MRI USE SCENARIO

To define the functional unit, the use scenario must first be defined. The use scenario includes applicable use modes, typical customer applications, and equipment capability. Use modes in view of the measurement of the energy consumption are defined as:

**Off mode:** The system functions into the minimum energy consumption state that the typical user can access e.g. through selection of off or shutdown, at the operator console.

**Ready-to-scan mode:** This mode represents the state of the system during patient handling and/or data evaluation and archiving, between individual scans.

**Scan mode:** The MRI is actively scanning the patient to generate images by sending high frequency waves and gradient pulses and reading the resulting variations in the magnetic field. The computing system interprets the data and generates the images.

To determine the time an MRI system remains in each mode, participants referenced confidential field usage records and estimated average values that could represent daily usage of MRI.

---

25 The low power mode defined in the MRI measurement methodology is not reported here as its power consumption has been assumed equal to ready-to-scan mode.
To evaluate the energy consumption, the most commonly used examinations were estimated by application specialists. Such values are also supported by external studies such as the “2007 MRI Market Summary Report”, May 2008, IMV Medical Information Division\textsuperscript{[26]}. This mix served as the "standard application mix" on which basis specific MRI protocols were defined and performed. Members agreed to use the top 5, normalized to 100% as shown in table 10.

**Table 10: Scan Mode application mix**

<table>
<thead>
<tr>
<th>Diagnostic Application</th>
<th>Normalized Distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Head</td>
<td>23,8%</td>
</tr>
<tr>
<td>Spine</td>
<td>24,8%</td>
</tr>
<tr>
<td>Abdomen</td>
<td>23,8%</td>
</tr>
<tr>
<td>Knee</td>
<td>19%</td>
</tr>
<tr>
<td>Angio</td>
<td>8,6%</td>
</tr>
</tbody>
</table>

According to companies’ experts the following daily usage has been defined:

- **Off:** 12h (Off mode)
- **Scan, ready-to-scan:** 10h (Ready-to-scan + scan mode)
- **Low power:** 2h (Reduced power consumption)

### 3.7.5. PATIENTS PER DAY

A very important feature of MRI is the patient/day ratio. The patient/day ratio measures the maximum number of patients (or examinations) that a MRI machine could scan in one day according to the examination distribution (use scenario) set as typical by the measurement methodology.

This value is determined by performing each examination (head, spine, abdomen, knee, angio) using phantoms\textsuperscript{[27]} but with real patient measurements (e.g. contrast agent injections, table moves, patient breath holds, etc). Using the distribution provided by the use scenario, it is possible to determine how many examinations could be performed in one day (how many patients could be examined per day).

The patient/day ratio is very important for at least 2 main reasons:

- The productivity of the machine represents high value information for the user (hospital/clinic).
- There is a linear correlation between the productivity and the energy consumption. MRI with higher patient/day ratio consumes more energy as shown by figure 7.

This means that MRIs with lower performances in terms of patients per day are usually consuming less energy. Reducing the number of patients per day could help reducing the energy consumption of MRI equipment. On the other hand, this is not acceptable as medical companies are committed to deliver equipment with improved performances/shorter examination times for patients.

As the technological evolution is moving towards machines with faster scan time and higher patient throughput (higher productivity) the energy consumption in absolute value could be expected to grow accordingly.

\textsuperscript{26} [www.imvinfo.com](http://www.imvinfo.com).

\textsuperscript{27} Phantoms are models made of plastic and fluids that simulate body parts and are used to test and calibrate MRI equipment.
3.7.6. **SIMPLIFIED MEASUREMENT METHODOLOGY**
As the Steering Committee decided to measure only the energy usage in off and ready-to-scan mode (see Appendix 3.7), the "Methodology for the measurement of the energy consumption of MRI" has been applied in a simplified version. Moreover, the power draw in low power mode as defined in methodology description has been assumed equal to the power draw in ready-to-scan mode.

3.7.7. **THE METHODOLOGY IN BRIEF**
The methodology requires and explains how to measure the following data needed for the SRI:

1. Power consumption in off mode
2. Power consumption in ready-to-scan mode
3. Duration of each one of the defined sequences

The duration of each examination is calculated as the sum of the time in scan mode (measured) and the time in ready-to-scan-mode (average value derived by companies’ experience and first simulations).

An evaluation spreadsheet calculates the following values:
1. Number of examinations per day: calculated from the duration of each examination and the examination distribution in the use scenario during 10 hours daily working time.
2. Energy consumption in off mode: calculated multiplying the power consumption in off mode by 12 hours
3. Energy consumption in ready-to-scan mode: the energy consumption of each examination is calculated multiplying the measured power consumption for the duration of ready-to-scan. The total energy consumption per day is obtained multiplying such values for the number of examinations per day.

All the details and procedures on how to measure the energy consumption are presented in the "Magnet Resonance Equipment (MRI) – Measurement of energy consumption“ document, available on the COCIR website.

3.7.8. **REQUIRED RESOURCES TO PERFORM THE MEASUREMENTS**
The measurement methodology requires the MRI to be available in a test lab. In alternative the test could be performed in a hospital or clinic environment.

The following tasks and technicians/specialists are required to measure one specific target MRI
equipment:

<table>
<thead>
<tr>
<th>TASK</th>
<th>TIME</th>
<th>Person</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compilation of the sequences</td>
<td>4h</td>
<td>Application specialist</td>
</tr>
<tr>
<td>Installation of the measurement tool</td>
<td>1h</td>
<td>Electrician</td>
</tr>
<tr>
<td>Preparation of the templates</td>
<td>1h</td>
<td>Specialist</td>
</tr>
<tr>
<td>Running the sequences</td>
<td>3h</td>
<td>Specialist, Measurement specialist</td>
</tr>
<tr>
<td>Measurement of Off mode and de-installation of measurement tool</td>
<td>1h</td>
<td>Specialist, Electrician, Measurement specialist</td>
</tr>
<tr>
<td>Data archiving</td>
<td>1h</td>
<td>Application specialist</td>
</tr>
<tr>
<td>Data evaluation</td>
<td>4h</td>
<td>Specialist</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>20h</strong></td>
<td></td>
</tr>
</tbody>
</table>

### 3.8. ECODESIGN TARGET FOR MRI

#### 3.8.1. SRI METHODOLOGY FOR ECODESIGN TARGET SETTING IN BRIEF

The fourth step of the SRI methodology sets the ecodesign target as the market average performance of the selected aspect, to be achieved in a time equal to the specific innovation cycle.

The SC Secretariat collects from companies the measurements of all MRI models and, on the basis of the reduction potentials determined by the PE INTERNATIONAL study on MRI, calculates the target scenarios:

- Baseline today
- Business as usual (BAU)
- Best not yet available technology (BnyAT)
- Beyond Business as usual

Based on these scenarios, the SRI SC decides on a feasible industry reduction target. Before it is approved, the industry target is proposed to the European Commission for discussion.

#### 3.8.2. MRI CATEGORIES

The SRI SC recognized that MRI equipment has different design intents, for specific clinical applications. The design intents result in energy consumption which is substantially different, due in large part to MR physics. For instance, a growing clinical need is for MRI systems with a large patient access (bore). Since MR physics is based on pulse sequences (switched magnetic field gradients and radio frequency pulses), the power needed for the pulse sequences increases as the diameter increases. Other features relevant to different image quality needs, such as number of data receiver channels, also affect energy consumption. It was recognized that a simple energy metric might cause confusion if systems with different clinical utilities are compared directly. As a result, member companies have developed a categorization table (see table 11).

---

28 For additional information on scenarios refer to SRIv2 documentation, Appendix V: www.cocir.org
### Table 11: MRI Equipment Categorization Table

| General information on categories included |  
|------------------------------------------|---
| - Matrix columns represent key differentiation characteristics that differentiate different clinical utilities of a system. |  
| - Each characteristic results in a designated amount of points and the total score of all characteristics will determine the overall category that a system belongs to |  
| **Key characteristics** |  
| **Field strength** |  
| 1.5T | 50 points  
| 3.0 T | 100 points  
| **Bore size** |  
| < 60 cm | 10 points  
| ≥ 60 & < 70 cm | 20 points  
| ≥ 70 cm | 30 points  
| **Maximum Gradient Amplitude per axis** |  
| < 35 mT/m | 40 points  
| ≥ 35 mT/m | 80 points  
| **Maximum Slewrate per axis** |  
| ≤ 100 mT/m/s | 20 points  
| ≥ 100 mT/m/s & < 150 mT/m/s | 30 points  
| ≥ 150 mT/m/s | 40 points  
| **Patient table** |  
| fixed table | 10 points  
| mobile table | 20 points  
| **Maximum channels** |  
| ≤ 16 channels | 15 points  
| ≥ 16 channels & < 64 channels | 35 points  
| ≥ 64 channels | 45 points  
| **Useable FOV cm²** |  
| ≤ 40 cm | 25 points  
| ≥ 40 & < 50 cm | 35 points  
| ≥ 50 cm | 45 points  
| **Total points** |  
| Clinical model - Category A | < 220 points  
| Hospital model - Category B | ≥ 220 & < 315 points  
| Research model - Category C | ≥ 315 points  

The MRI units sold in 2010 and 2011 are reported in figure 8 in percentage according to the 3 identified categories.

Figure 8: MRI – Distribution of units sold* in 2010 and 2011 in EU

The required high level performances involve higher energy consumption, due to the high magnetic and gradient field performance, number of receiving channels, bore and field-of-view size. For this reason, the potential for improvement is extremely limited and should be investigated with extreme care to avoid that possible technical solutions to reduce the energy consumption (adopted for category B equipment) could compromise the innovation potential.

For the above mentioned reasons the SRI Steering Committee decided not to set targets for such equipment and to evaluate the feasibility of reducing the energy consumption without compromising performances and benefits for patients.

3.8.4. FUNCTIONAL UNIT

30 EU 27 market data provided by companies for each model placed on the market in calendar year 2010 and 2011. As only category B products have been included in the SRI scope, sales data on category A and C are not collected every year.
The functional unit is the reference ensuring the comparability of power consumption of different products and their developments over time.

As identified by the study on improvement potential for MRI, the functional unit for MRI is the number of patients that can be examined per day. Such number, as already presented, is not fixed a priori but depends on the hospital workflow, the administrative time, the nature of examinations, the required quality and functionality and furthermore the power and performance of the machine. It is determined measuring the duration of each examination (scan time: measured + ready-to-scan time: set) and applying the examination distribution to the 10 hours working time of the machine.

3.8.5. **METRIC – ENERGY CONSUMPTION IN OFF AND READY-TO-SCAN ONLY**

The energy consumption of MRI is the sum of the energy consumption in the three different modes (off, ready-to-scan and scan).

The initial measurements run on 12 models and the results of the study on MRI potential for improvement have shown that:

1. Measuring the energy consumption in scan mode is complex, expensive and time consuming, as examined in Appendix II.I.

2. The potential for reducing the energy used to perform the scan is limited due to the physics of the process. A certain amount of energy is needed to stimulate the response from the body and to be read by receivers.

3. Improvements in scan mode could be achieved by defining new technologies that use different sequences with less energy needed. Such improvements could not be recorded by the methodology at the moment, as the sequences are set. Not setting the sequences would render difficult to compare the measurements as it changes the functional unit.

Therefore, the SRI Steering Committee decided to not consider the energy in scan mode for the determination of the ecodesign target.

The adopted metric for setting the target for MRI is the energy usage per model per day (kWh/unit day) in off and ready-to-scan mode to perform a certain number of examinations according to the use scenario.

The target is to be expressed as the average daily consumption per model in off and ready-to-scan mode:

\[ \text{kWh} \text{(off, ready-to-scan)/unit day} \]

This choice reflects the part of the energy consumption that could be reduced by ecodesign programs and takes into account the productivity of the MRI as the time in ready-to-scan mode is not defined but varies. In fact, even if the ready-to-scan time is defined per examination, the number of examinations per day depends on the total examination time, which account also for the scan time.

3.8.6. **INNOVATION CYCLE**

The innovation cycle is defined as the time needed to develop new or enhanced products and place them on the market. For medical devices it could vary from 3 years to 7, depending on the complexity of the innovation being brought to market.
The below listed activities for MRI requires:

- Research and development: 1 year
- Realization, Verification and Validation: 3 years
- Regulatory Approvals: 1 year

The innovation cycle for MRI therefore corresponds to 5 years.

3.8.7. SETTING THE ECODESIGN TARGET

The SRI methodology for target setting has been developed on the base of the experience gathered with the pilot ultrasound project. In particular the Business-as-usual scenario (BAU) is based on the assumption that the energy consumption of the modality under consideration will get lower year after year due to existing ecodesign programs and due to the improvements of other technologies according to implementing regulations under the Ecodesign Directive or voluntary measures. This assumption has proven true for the ultrasound pilot project.

The PE INTERNATIONAL study on MRI shows that this assumption is not true for MRI. New functionalities, larger bore diameters, increased magnetic field strength and more powerful gradient and RF amplifiers are going to increase the energy demand to meet clinical needs of medical care.

Therefore the BAU scenario, defined under the assumption that all companies will reach the front runner today at the end of the innovation cycle, has been redefined for MRI according to the findings of the PE INTERNATIONAL study on MRI improvement potentials and used as the baseline.

According to the findings of PE INTERNATIONAL, the BAU baseline shows an increase in the energy demand which can be mitigated by the reduction of energy usage in the most favorable case (BnyAT) where all possible improvements are implemented at the same time by all companies (extreme assumption not in line with technological limits). Therefore the BnyAT scenario should be re-defined accordingly as the result of the application of the companies' potentials to the newly defined BAU baseline.

**BAU Scenario**

PE International estimated in the BAU scenario an increase in energy consumption (off+ready-to-scan) of 16.68% assuming that by 2017, half of the category B product sold on the market will have an energy consumption comparable to the energy consumption of Category C products today.

This estimation has been reviewed later with the availability of additional measurement data and estimations provided by each Company of its own BAU scenario according to the specific corporate strategies.

An increase around 12% in the energy consumption by 2017 has been considered an assumption better reflecting the current trends.

**Beyond BAU scenario and correction factors**

According to the experience gathered with the ultrasound Pilot project the SRI methodology assumes that the front runner is the Company with the lowest potential for improvement. The study on Improvement Potentials coupled with measurement data showed for MRI a different situation. The front runner estimated an improvement potential that is quite high compared to other companies.

---

31 The new methodologies for the scenarios definition (BAU and BnyAT) have been included in the SRIv3 revision which has been published with this Report in 2013.
This can be interpreted as the result of extensive research in ecodesign that allows the front runner to foresee the application of technical solutions that are not evident to other companies to improve the energy performances. This represents an important example of how ecodesign could drive innovation.

Applying the SRI methodology under this circumstances is not possible, otherwise the industry target will result even higher than what has been estimated as the highest possible improvement.

The SRI Steering Committee decided to use correction factors applied to the individual company maximum improvement to derive the company targets and the Industry targets (weighted average against sales). It has been assumed that companies could achieve 75% of the maximum possible improvement and 50% for the front runner to take into consideration the higher marginal costs.

**Scenarios**

The four scenarios have been redefined accordingly as:

- **Baseline today**
- **Business as usual scenario (BAU) according to the SRIv2 methodology**, used as reference value and showing the fleet performance of the front runner.
- **Business as usual (BAU) according to the SRIv3 methodology**: scenario for year 2017 where the average daily energy usage per model is expected to be increased around 12% compared to baseline today.
- **Best not yet available technology (BnyAT)**: scenario for year 2017 where the average daily energy usage per model is expected to decrease around 5,4% compared to the baseline 2011.
- **Beyond Business as usual**: scenario for year 2017 derived applying correction factors to companies BnyAT where the application of the SRI will compensate the increase in energy consumption due to added functionalities maintaining the energy consumption constant (0,73% decrease compared to the 2011 baseline).

The maximum possible reduction potential identified for each Company is used to calculate the average value, **15,63%**, that is used to define the Best-not-yet-available scenario. The PE International study collected the individual company data that cannot be disclosed due to confidentiality reasons.

The four scenarios calculated on all 14 measured models in category B are indicated in the following table:

<table>
<thead>
<tr>
<th>Scenario (kWh/unit day)</th>
<th>Company</th>
<th>Average daily consumption in off and ready-to-scan per unit (kWh/d)</th>
<th>Range for setting targets compared to baseline 2011</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline 2011 (kWh/d)</td>
<td>XX XX XX XX XX</td>
<td>227,4</td>
<td></td>
</tr>
<tr>
<td>BAU 2017 according to SRI methodology</td>
<td>XX XX XX XX XX</td>
<td>176</td>
<td>Front runner fleet performance</td>
</tr>
<tr>
<td>BAU 2017 (kWh/d)</td>
<td>XX XX XX XX XX</td>
<td>254,9</td>
<td>+12,07%</td>
</tr>
<tr>
<td>BnyAT 2017 (kWh/d)</td>
<td>XX XX XX XX XX</td>
<td>215,2</td>
<td>-5,38%</td>
</tr>
<tr>
<td>Beyond BAU 2017 (kWh/d)</td>
<td>XX XX XX XX XX</td>
<td>225,8</td>
<td>-0,73%</td>
</tr>
</tbody>
</table>

Grey cells: confidential data
The baseline scenario is obtained as the weighted average of the energy performance of all models in kWh\textsubscript{(off+ready-to-scan)/unit\textperiodcentered day} against the sales\textsuperscript{32}.

The result means that companies producing MRI equipment participating in the SRI commit not to increase the energy consumption in off and ready-to-scan mode of the average model in 2017 compared to the 2011 baseline. If the SRI was not in place, the energy consumption would have increased around 12\% by 2017.

Table 12: Calculated values for year 2010-2011\textsuperscript{33} and forecast until 2017 under the assumption of a linear trend.

<table>
<thead>
<tr>
<th>Year</th>
<th>Sold units\textsuperscript{34}</th>
<th>Total daily energy consumption (kWh)\textsuperscript{38}</th>
<th>Average daily energy consumption per unit (kWh/d)</th>
<th>Beyond BAU</th>
<th>BAU</th>
</tr>
</thead>
<tbody>
<tr>
<td>2011</td>
<td>✓394</td>
<td>✓89.011</td>
<td>✓225,92</td>
<td>225,61</td>
<td>230,39</td>
</tr>
<tr>
<td>2012</td>
<td></td>
<td></td>
<td></td>
<td>225,30</td>
<td>234,87</td>
</tr>
<tr>
<td>2013</td>
<td></td>
<td></td>
<td></td>
<td>225,00</td>
<td>239,34</td>
</tr>
<tr>
<td>2014</td>
<td></td>
<td></td>
<td></td>
<td>224,69</td>
<td>243,81</td>
</tr>
<tr>
<td>2015</td>
<td></td>
<td></td>
<td></td>
<td>224,38</td>
<td>248,29</td>
</tr>
<tr>
<td>2016</td>
<td></td>
<td></td>
<td></td>
<td>224,07</td>
<td>252,76</td>
</tr>
<tr>
<td>2017</td>
<td></td>
<td></td>
<td></td>
<td>224,07</td>
<td>252,76</td>
</tr>
</tbody>
</table>

Figure 9: MRI target scenarios

\textsuperscript{32} This value differs slightly from the baseline value of the PE INTERNATIONAL study as the study identified the improvement potential of a representative model therefore considering a simple average of the measured models. Therefore the data presented in this report better reflects market reality.

\textsuperscript{33} Values are slightly different from the ones in SRI Report 2012 due to a retrospective re-categorization of products by one company causing a variation of SRI sales and related energy consumption data of 0,15\% in 2011 and 0,13\% in 2012 compared to the values reported in previous SRI Status Reports.

\textsuperscript{34} Sold units data provided by companies for each model placed on the market in the calendar year 2011.

\textsuperscript{35} Measured energy consumption data provided by companies for each model placed on the market in the calendar year 2011.
3.8.8. COMPANY TARGETS
According to the SRI methodology, each member company adopts an internal company target which enables achievement of the industry target.
Every year the SRI SC Secretariat can evaluate the achievement of each company by comparing the baseline with the measured average performance of all models from each company placed on the market each year.
As the improvement is not a linear process only at the end of the 5 years period it would be possible to evaluate whether the Company targets have been achieved or not.

Member company targets are confidential unless a company wishes to disclose its own.

3.9. RELATIONSHIP BETWEEN SCAN AND READY-TO-SCAN KWH
Figure 10 represents the relationship between the energy use in scan mode and the energy use in ready-to-scan mode measured on 12 models.
The linear regression shows a good correlation ($R^2=0.77$) which allows to determine the total daily consumption of a MRI given the consumption in off and ready to scan mode.

\[
\text{Scan kWh/d} = 0.7688 \times \text{Ready-to-scan kWh/d} + 4.835
\]

The relationship is valid for year 2011 and will gradually change in time due to the different trends in energy usage in the different modes.
APPENDIX IV

4. COMPUTED TOMOGRAPHY

4.1. GENERAL DESCRIPTION OF COMPUTED TOMOGRAPHY

Computed tomography is a medical imaging procedure that utilizes computer-processed X-rays to produce tomographic images or 'slices' of specific areas of the body. CT is used in medicine as a diagnostic tool and as a guide for interventional procedures. Sometimes contrast materials, such as intravenous iodinated contrast, are used. This is useful to highlight structures such as blood vessels that otherwise would be difficult to delineate from their surroundings. Using contrast material can also help to obtain functional information about tissues.

X-ray slice data is generated using an X-ray source that rotates around the object; X-ray sensors are positioned on the opposite side of the circle from the X-ray source.

Since the nineties CT scanners can process not only individual cross sections but continuously changing cross sections as the gantry, with the object to be imaged slowly and smoothly slid through the X-ray circle. These are called helical or spiral CT machines. Their computer systems integrate the data of the moving individual slices to generate three-dimensional volumetric information (3D-CT scan), in turn viewable from multiple different perspectives on attached CT workstation monitors. This type of data acquisition requires enormous processing power, as the data are arriving in a continuous stream and must be processed in real-time.

Detectors

The earliest sensors were scintillation detectors, with photomultiplier tubes excited by (typically) cesium iodide crystals. Cesium iodide was replaced during the 1980s by ion chambers containing high-pressure Xenon gas. These systems were in turn replaced by scintillation systems based on photodiodes instead of photomultipliers and modern scintillation materials with more desirable characteristics.

To be able to obtain a good quality image, detectors have to reach a specific temperature (steady state) from the off state. This process could take different times according to the specific CT and detector technology.

Reconstruction engines

Once the scan data has been acquired, the data must be processed using a form of tomographic reconstruction, which produces a series of cross-sectional images. In terms of mathematics, the raw data acquired by the scanner consists of multiple "projections" of the object being scanned. These projections are effectively the Radon transformation of the structure of the object. Reconstruction essentially involves solving the inverse Radon transformation.

Recently, manufacturers have developed iterative physical model-based maximum likelihood expectation maximization techniques. These techniques are advantageous because they use an internal model of the scanner's physical properties and of the physical laws of X-ray interactions. Iterative techniques provide images with improved resolution, reduced noise and fewer artifacts, as well as the ability to greatly reduce the radiation dose in certain circumstances. The disadvantage is a very high computational requirement, but advances in computer technology and high-performance computing techniques, such as use of highly parallel GPU algorithms, now allow practical use.

https://www.medicalradiation.com
4.2. MARKET DATA

The ✓537 ✓ companies participating in the SRI for the CT sector represent a total turnover in Europe of ✓46238 million euros in 2016 and cover about 100%39 of the European market.

Table 13: CT - EU40 market data

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Computed tomography (CT)</td>
<td>✓402 M€</td>
<td>✓489 M€</td>
<td>✓462 M€</td>
<td>100%</td>
</tr>
</tbody>
</table>

4.3. IDENTIFICATION OF THE MOST SIGNIFICANT ENVIRONMENTAL ASPECT FOR CT

Computed tomography equipment has resulted as the second modality to be targeted in 2012 on the base of Step 1 and Step 2 of the methodology as shown in table 5.

According to Step 3 of the methodology as summarized in chapter 3, the data provided by companies are used to rank the different environmental aspects. Table 14 shows that energy consumption during the product lifecycle use phase has been identified as the top environmental aspect, representing around 75% of the impacts on the life cycle of a CT equipment.

Table 14: Identification of most significant environmental aspect

<table>
<thead>
<tr>
<th>Identification of most significant environmental aspect</th>
<th>Average internal ranking</th>
<th>% of total life-cycle</th>
<th>Final COCIR Ranking</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy use</td>
<td>1</td>
<td>75%</td>
<td>1</td>
</tr>
<tr>
<td>Non-ferrous metals and alloys</td>
<td>2</td>
<td>11%</td>
<td>2</td>
</tr>
<tr>
<td>Ferrous metals and alloys</td>
<td>3</td>
<td>6%</td>
<td>3</td>
</tr>
</tbody>
</table>

The expert judgment provided by companies’ experts shows that there is a potential for improvement, even if limited, in the energy consumption of CT. Risks have been identified regarding the impact on patient throughput and innovation. In particular extreme care should be used to ensure that the energy usage reduction is not going to affect the radiation dose or the development towards lower dosage in the future.

37 GE Healthcare, Hitachi Medical Systems Europe, Philips Healthcare, Siemens Healthcare, Toshiba Medical Systems Europe
38 COCIR Imaging Market Statistics source (SHARE).
39 COCIR estimation based on 2015 SHARE data
40 Countries included: Estonia, Latvia, Lithuania, Bosnia, Bulgaria, Croatia, Czech Republic, Hungary, Poland, Romania, Serbia, Slovakia, Slovenia, Ukraine, Portugal, Spain, Denmark, Finland, Norway, Sweden, Ireland, UK, Austria, Belgium, France, Germany, Greece, Italy, Netherlands, Switzerland, Albania, Macedonia.
4.4. MODES
Four functioning modes have been defined for CT equipment.

**Off:** The system is shut down, according to the user manual. The system consumes no energy.

**Low-power:** The system functions into the minimum energy consumption state that the user can select according to the user manual.

**Idle mode:** This mode represents a state of the system when fully powered but no scan has been prescribed. This mode does **NOT** include x-ray tube rotor or gantry rotation.

**Scan mode:** This mode represents the state of the system between individual scans and during scans (e.g. during patient handling, examination planning, contrast agent injection and active scanning with x-ray generation). This mode includes tube rotor rotation, gantry rotation and generation of image.

The power consumption of CT in the four modes is represented in figure 11 for the typical use scenario used by the SRI. Figure 12 provides a detailed overview of power draw during scan.

*Figure 11: Power usage of a CT scanner over a typical day in hospital. The Idle mode has been simplified for illustration purposes. The 45 minutes at the end of the 12 working hours includes 35 minutes of the last scan and 10 minutes in Idle mode.*
4.5. MODULARIZATION

The most important power consuming modules of the entire CT system have been identified\textsuperscript{41}.

<table>
<thead>
<tr>
<th><strong>Tube and generator chain:</strong></th>
<th>X-ray tube and all the power supplies used to power the tube and all connected devices.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Detector:</strong></td>
<td>Device reading the x-ray radiation and converting it into digital signal</td>
</tr>
<tr>
<td><strong>Power distribution unit and other power supplies</strong></td>
<td>Subsystems required for electrical energy conversion and distribution</td>
</tr>
<tr>
<td><strong>Computation, Controls:</strong></td>
<td>Units dedicated to the reconstruction of the CT image and the control of the CT system</td>
</tr>
<tr>
<td><strong>Cooling:</strong></td>
<td>A collection of subsystems required for thermal management of CT components</td>
</tr>
<tr>
<td><strong>Patient table:</strong></td>
<td>Patient table with electric movement controls</td>
</tr>
<tr>
<td><strong>Gantry Motor:</strong></td>
<td>Electric motor turning the gantry around the patient during scan time.</td>
</tr>
</tbody>
</table>

\textsuperscript{41} PE International: Computer Tomography (CT) - Study on the potential for environmental improvement by the aspect of energy efficiency
4.6. MEASUREMENT OF ENERGY CONSUMPTION

4.6.1. HISTORY

The SRI Steering Committee started in October 2010 to develop a methodology to measure the energy consumption of CT equipment, as there are no available standards that could be used.

The methodology ensures that:
- Results of the measurements are comparable and repeatable
- The measured energy consumption has a strict relation with the real energy consumption in everyday hospital use. Such information is important for hospitals and clinics to understand the real running cost of CT to plan and allocate correctly their budget
- The procedure does not involve disproportionate costs or resources
- The measurement results allow the determination of all the information that could be considered relevant for the SRI target setting process but could also be useful for CT users.

The methodology is available for download at COCIR website.

4.6.2. THE MEASUREMENT METHODOLOGY IN BRIEF

The SRI SC decided to use as functional unit a typical day of functioning of the CT scanner in hospital environment. The SRI SC determined as a realistic use scenario representing a typical day in the hospital 20 examinations over 12 hours.

To determine representative CT clinical procedures, the study “European Guidance on Estimating Population Doses from Medical X-Ray Procedures”, published by Directorate-General for Energy and Transport in 2008, is used here to determine the distribution of procedure types. From this report the defined procedure distribution used within is:

<table>
<thead>
<tr>
<th>Examination</th>
<th>Distribution</th>
<th>Examinations per day</th>
</tr>
</thead>
<tbody>
<tr>
<td>CT head</td>
<td>42,9%</td>
<td>9</td>
</tr>
<tr>
<td>CT chest</td>
<td>15,9%</td>
<td>3</td>
</tr>
<tr>
<td>CT spine</td>
<td>16,9%</td>
<td>3</td>
</tr>
<tr>
<td>CT abdomen</td>
<td>24,2%</td>
<td>5</td>
</tr>
</tbody>
</table>

Second, within each procedure type, the specific scan parameters representing each exam type have been chosen to harmonize across manufacturers to general values:

The 12 hours have been subdivided in 20 intervals of 35 minutes each (as shown in table 15).

**Table 15: Allocation of body regions scans during the 12 hours working time**

<table>
<thead>
<tr>
<th>Type of examination</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
<th>14</th>
<th>15</th>
<th>16</th>
<th>17</th>
<th>18</th>
<th>19</th>
<th>20</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H</td>
<td>A</td>
<td>H</td>
<td>T</td>
<td>H</td>
<td>A</td>
<td>S</td>
<td>H</td>
<td>T</td>
<td>H</td>
<td>A</td>
<td>S</td>
<td>H</td>
<td>T</td>
<td>H</td>
<td>A</td>
<td>S</td>
<td>H</td>
<td>A</td>
<td>H</td>
<td></td>
</tr>
<tr>
<td>Duration (min)</td>
<td>10</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>10</td>
<td>720</td>
</tr>
</tbody>
</table>

Trial executions of testing showed a further simplification of the test procedure is possible by approximating the range of protocols by only the Abdomen scan. Table 16 shows the allocation...
of time during the working hours.

**Table 16: Allocation of abdomen scans during the 12 hours working time**

| Type of examination | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | Total |
| Duration (min)      | 10 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 35 | 10 | 720 |

\[ E_{\text{scan}} = 20 \times E_{\text{Abdomen}} \]

Each interval of 35 minutes includes (see figure 13) time in Scan mode, typical of each CT scanner and the remaining time in Idle mode. The scan interval, as shown in figure 13, is the time between the scan prescription and the moment the power usage is back to Idle mode.

![Figure 13: Representation of power draw of a CT scanner during LowPower, Idle and Scan mode](image)

The energy consumption of CT can be calculated by summing the measured energy consumption in each mode, where applicable, to a given use scenario:

\[
\text{Energy use} = E_{\text{off}} + E_{\text{idle}} + n \times E_{\text{scan}} + E_{\text{low}}
\]

### 4.6.3. ASPECTS OF THE METHODOLOGY TO BE IMPROVED

The measurement methodology is a powerful tool that allows the measurement of energy consumption based on a use scenario that is very close to everyday practice.

Nonetheless the methodology still has some weak points that the SRI Steering Committee is committed to improve in the coming years.

**Benefit for patients**

In its current form the methodology takes marginally into account the benefits for patients. Companies are working to provide better technologies with improved functions, able to provide better comfort and benefits for patients such as:

- Image quality and resolution
- Integration with other technologies
- Shorter exam durations
- Noise insulation systems
- Alternatives to the use of contrast agents
- Dose reduction

Most of those options require higher energy use. In particular the reduction of the dose is one of the most researched issues at the moment. The reduction of the x-ray dose involves more
powerful and sophisticated reconstruction capabilities, which translates into higher computational power and therefore higher energy consumption. An increase in the average energy consumption for CT due to the increased functionality and/or dose reduction is not recorded by the methodology.

4.7. ENERGY CONSUMPTION

SRI participating companies measured most of the models placed on the market in 2012. The average energy usages are reported in table 17 for the 4 functioning modes.

Table 17: Average CT energy consumption in different modes in case the CT is switched to LowPower mode overnight.

<table>
<thead>
<tr>
<th>Mode</th>
<th>Typical time in mode per day</th>
<th>Average energy consumption per day</th>
<th>Estimate of % energy in use phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off mode</td>
<td>0</td>
<td>0</td>
<td>0%</td>
</tr>
<tr>
<td>LowPower mode</td>
<td>12</td>
<td>12,2</td>
<td>25%</td>
</tr>
<tr>
<td>Idle mode</td>
<td>10,8</td>
<td>31,1</td>
<td>62%</td>
</tr>
<tr>
<td>Scan mode</td>
<td>1,2</td>
<td>6,1</td>
<td>13%</td>
</tr>
<tr>
<td>Total</td>
<td>50</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The daily energy consumption depends on which mode is selected for 12h night time.

Table 18: Average CT energy consumption in different scenarios.

<table>
<thead>
<tr>
<th>Scenario</th>
<th>Description</th>
<th>Average energy consumption per day kWh/d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Off</td>
<td>The CT is switched to Off mode during the 12h night time. No energy consumption.</td>
<td>37</td>
</tr>
<tr>
<td>LowPower</td>
<td>The CT is switched to LowPower mode during the 12h night time</td>
<td>50</td>
</tr>
<tr>
<td>Idle</td>
<td>The CT works in Idle mode during the 12h night time</td>
<td>72</td>
</tr>
</tbody>
</table>

According to the modularization defined in chapter I.V, the energy consumption can be allocated in the different modules and modes as indicated in figure 14.

\[42\text{ Values taken from the PE International Study : Computed Tomography (CT) - Study on the potential for environmental improvement by the aspect of energy efficiency – Page 23} \]

\[43\text{ Percentages are calculated against a scenario assuming the scanner working in the corresponding mode for the 12h night time. For this reason percentages cannot be summed to obtain 100%} \]

\[44\text{ Values taken from the PE International Study : Computed Tomography (CT) - Study on the potential for environmental improvement by the aspect of energy efficiency – Page 29} \]
Figure 14: Energy consumption over a typical day in LowPower, Idle and Scan mode allocated to the different modules of the CT scanner⁴⁵.

4.8. REDUCTION OF X-RAY DOSE – COCIR VOLUNTARY COMMITMENT WITH HERCA

Media reports, the public and governmental authorities have placed ionizing radiation exposure and dose reduction measures in medical imaging high on the public health agenda. This increases the awareness among the various stakeholders, such as clinical professionals, equipment manufacturers, regulators, hospital managers, patients, etc. New requirements and the implementation of future workflow concepts on dose management and dose reporting are currently being considered around the world. Since February 2010, regular discussions are taking place between COCIR and the Heads of European Radiation Competent Authorities (HERCA) requesting our industry to commit in reducing radiation dose for CT equipment. As the developers of sophisticated scanners, CT manufacturers acknowledge their unique role in the process to help optimize patient CT dose in the health care setting.

A dedicated COCIR Task Force was created to respond to HERCA’s request and a COCIR CT manufacturers’ voluntary commitment was released in May 2011.

The aim of this commitment is to further the initiatives of improving dose reporting, promoting transparency in dose efficacy, continuing reduction of medical exposures, and provision of specific training curricula.

The manufacturers agree to complete the voluntary commitments outlined within and provide yearly updates on:
1. Characterization of CT Systems Standardized Benchmarking
2. Implementation of dose reduction measures in CT
3. Dose management and reporting
4. Provision of specific training curricula

COCIR CT manufacturers have been developing and providing dose reduction features on CT systems for many years, and this trend continues today:

⁴⁵ PE International: “Computer Tomography: Study on the potential for environmental improvement by the aspect of energy efficiency”, page 23.
- Patient Protocol Selection Guidance
- Automatic Tube Current Modulation (ATCM) and X-ray Initiation
- Precise X-ray Field shaping
- Dose Efficient Design
- Dose reporting and Awareness
- Training Opportunities
- Pediatric Protocols
- Dedicated Infant Imaging Mode
- Advanced Tube and Collimator Design
- Dose Efficient Detection
- Dose Display and Recording
- Optimized Image Reconstruction

Some of the above mentioned technological solution could bring to higher energy consumption while trying to reduce dose. Their mutual feedbacks and interactions on energy usage are extremely hard to forecast. For this reason, companies could not provide an estimation of the evolution of energy usage for the next years, but an increase in idle energy is most likely.

All the documentation related to the COCIR Voluntary Commitment with HERCA is available on the COCIR website.

4.9. USER BEHAVIOUR
Given the energy consumption data in the two scenarios in the previous chapter, the SRI SC looked carefully into a fundamental issue influencing energy consumption patterns: user behavior.

The analysis has been performed on two levels:
1. Company field data
2. Literature

CT scanners used in emergency care settings are rarely turned Off or to Low Power mode during night hours, but are kept in idle mode for rapid imaging to meet the needs of critical care. The SRI SC confirmed that many scanners used in other radiology settings are left in an idle mode during overnight hours. They are rarely turned Off or to LowPower mode, in fact are typically powered down only for maintenance or software reboot.

The analysis confirmed that around 90% of the scans are performed between 8.00 am and 8.00 pm (12h working time per day), but also that up to 70% of the scanners are not switched to Off/LowPower mode overnight. On the average, the remaining 30% is switched off only 50% of the time.

Considering the reduced energy consumption over 12 hours of the Off mode (0 kWh) or LowPower mode (12,3 kWh) compared to idle mode (34,6 kWh), a correct user behavior could allow to save from 12,23 to 34,6 kWh per day just by switching the CT scanner from idle. This value represents up to 48% of the energy used daily.

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6 IEEE Explore: Energy consumption of VA hospital CT scans / Study (Denmark) - Energy Efficiency in Hospitals and Laboratories
4.10. IMPROVEMENT POTENTIAL
The PE International study\(^\text{47}\) on the improvement potentials regarding the energy consumption for CT shows that:

1. **No improvements possible in scan mode:** considering the short duration of scans and the priority given to the reduction of the dose for patients, no improvements in the energy consumption are achievable. As a consequence of improvements in other modes, a small increase in scan energy is possible.

2. **Limited improvements in idle mode:** Considering the short periods of idle mode in between scans and the need for the user to have the scanner always ready to scan for the next examination, reduction of energy use during idle mode is hardly possible. Switched off components would require time to be reactivated thus hampering availability of the scanner in a very short time.

3. **Significant improvement potential in LowPower mode:** LowPower mode offers the greatest potential for improvement. In LowPower mode modules are active to provide a fast reactivation of the CT scanner to idle mode.

![Figure 15: Maximum improvement potentials of CT technology per mode](image)

It is important to note that the reported improvements are the maximum improvement potentials, also taking into consideration technologies that are not yet available today (ByAT). Therefore any final target would be necessarily lower. It is also important to consider that the LowPower mode accounts for just a 24,5% of the total daily energy consumption and therefore the 53% improvement even if possible, will end up in a 13% reduction.

The analysis of user behavior showed that about 70% of CT scanners are never switched to Off/LowPower during the 12h night time and the remaining 30% only 50% of the time on the average. This means that, unless user behavior can be influenced, any reduction of the device energy usage in LowPower mode may have a limited net effect for users (around 1,9%), since the LowPower mode is rarely used.

\(^{47}\) PE International: “Computer Tomography: Study on the potential for environmental improvement by the aspect of energy efficiency”, page 31. Available at www.cocir.org
4.11. APPLICATION OF SRIV3 METHODOLOGY TO CT

4.11.1. SRI METHODOLOGY FOR ECODESIGN TARGET SETTING IN BRIEF

The fourth step of the SRI methodology sets the ecodesign target as the market average performance of the selected aspect, to be achieved in a time equal to the specific innovation cycle.

The SC Secretariat collects from companies the measurements of all CT models and, on the basis of the reduction potentials determined by the PE INTERNATIONAL study on CT, calculates the target scenarios:

• Baseline today
• Business as usual (BAU)
• Best not yet available technology (BnyAT)
• Beyond Business as usual.

Based on these scenarios, the SRI SC decides on a feasible industry reduction target. Before it is integrated into company design targets, the industry target is proposed to the European Commission for discussion.

The results of this step are two types of targets:

• Industry target: that’s the target that all the participating companies have to achieve as the average of the market and is equal (unless a different decision is justified) to the value provided in the Beyond as usual scenario. This target is the target against which the success of the initiative has to be assessed.

• Individual company targets: Those are improvement targets that each company can derive from the reported scenarios. A company absolute target is equal to the average value provided by the BnYAT scenario. Such targets are used as an internal tool to keep track of improvements, to decide corrective actions and to ensure companies’ commitment.

4.11.2. SCOPE

The SRI Steering Committee decided to exclude certain specific CT models from the SRI methodology. In particular:

– 256 slices (or higher) scanners
– Dual source scanners

Dual source scanners are produced only by one Company today. Those are top level scanners equipped with two x-ray tubes. Both dual source and 256 (or higher) slice scanners represent high-end models, with increased functionality and top performances which involve higher energy consumption. Therefore the potential for improvement is extremely limited and should be investigated with extreme care to avoid that possible technical solution to reduce the energy consumption could compromise or reduce the performances.

4.11.3. FUNCTIONAL UNIT

The functional unit is the reference ensuring the comparability of power consumption of different products and their developments over time.

As identified in the CT energy measurement methodology, the functional unit is a typical daily usage in hospital environment.

48 For additional information on scenarios refer to SRIV3 documentation, Appendix V: www.cocir.org
4.11.4. INNOVATION CYCLE
The innovation cycle is defined as the time needed to develop new or enhanced products and place them on the market. For medical devices it could vary from 3 years to 7, depending on the complexity of the innovation being brought to market.

The below listed activities for CT requires:

- Research and development - 1 year
- Realization, Verification and Validation - 3 years
- Regulatory Approvals - 1 year

The innovation cycle for CT therefore corresponds to 5 years.

4.11.5. SETTING THE ECODESIGN TARGET
The analysis and studies performed by the SRI SC and PE International revealed so far two important aspects of CT technology that have to be taken into account:

1. The potential for improvement is limited: improvements can be achieved mainly in LowPower mode, which contribution to the daily energy consumption is limited (around 26% of daily energy usage) and rarely used (about 70% of all scanners are not switched to LowPower during 12h night time).

2. Ongoing activities and voluntary commitment of COCIR companies for the reduction of the x-ray dose are going to affect the energy consumption in scan and idle mode. While it is difficult to forecast the interaction of technical solutions and their effects on energy consumption, it is very likely that the energy consumption in idle will increase due to higher computation power required by more powerful iterative image reconstruction engines and longer computational time.

3. The energy savings allowed by the Off mode or LowPower mode, already implemented on CT scanners today, are not achieved due to user behavior. Any improvement achieved with technical solutions in such modes would not be used anyway.

4.11.6. INFLUENCING USER BEHAVIOUR
As shown by the PE International study, already existing Off and LowPower modes could ensure an energy saving between 30% and 45% of daily energy consumption. Nonetheless such options are not used by users.

Any technical solution to reduce the energy consumption could only improve the usage in LowPower mode and hardly in Idle mode.

In particular during working hours CT scanners work in Idle mode in-between scans, therefore for a quite short period. Reducing energy consumption in such periods would be complex as reactivation to idle mode could take quite some time which could be non-compatible with clinical needs.

The SRI SC concluded that the greatest reduction in energy usage can be achieved by influencing the users’ behavior through proper education and information about the possible energy savings related to an environmental friendly use.

Energy savings achievable by improving the use of LowPower mode can amount up to 8,8 MWh per year. Figure 16 shows the savings on a daily basis.
Figure 16: Energy consumption scenarios based on time spent by the scanner in LowPower mode during the 12h night time

Figure 17 shows the total daily energy consumption resulting from different duration of LowPower mode. The right column shows the energy usage in case the scanner is switched to Off mode during the 12h night period.

Figure 17: Energy savings achievable by using LowPower and Off mode during 12h night time

Ecodesign goals decided by the SRI SC to achieve energy savings by influencing user behavior towards an environmentally friendly use practice are reported in Chapter 6.

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49 PE International: “Computer Tomography: Study on the potential for environmental improvement by the aspect of energy efficiency”, page 29.
50 PE International: “Computer Tomography: Study on the potential for environmental improvement by the aspect of energy efficiency”, page 30.
APPENDIX V

5. X-RAY IMAGING EQUIPMENT

5.1. GENERAL DESCRIPTION OF X-RAY IMAGING EQUIPMENT

Radiography is an imaging technique that uses X-rays to view the internal structure of a non-uniformly composed and opaque object such as the human body. To create the image, a heterogeneous beam of X-rays is produced by an X-ray generator and is projected toward the patient’s body. A certain amount of X-ray is absorbed by the body, which is dependent on the particular density and composition of the area. The X-rays that pass through are captured by a detector (either photographic film or a digital detector). The detector can then provide a superimposed 2D representation of all the body's internal structures.

X-ray technologies

X-ray imaging comprises a lot of different applications for specific body parts or clinical needs. X-ray can be divided into two main groups: Fluoroscopy and Radiology.

Fluoroscopy is an imaging technique that uses X-rays to obtain real-time moving images of the internal structures of a patient through the use of a fluoroscope. In its simplest form, a fluoroscope consists of an X-ray source and fluorescent screen between which a patient is placed. However, modern fluoroscopes couple the screen to an X-ray image intensifier and CCD video camera allowing the images to be recorded and played on a monitor. The x-ray generation may last for several minutes at very low intensity.

Some forms of radiography include:
- Angiography
- Fluoroscopy
- Surgery

Radiography is used for fast, highly penetrating images, and is usually used in areas with high bone content. X-rays are generated for milliseconds at higher intensity.

Some forms of radiography include:
- Radiography
- Gen&Uro
- Mammo
Digital detectors and image intensifiers

Traditional X-ray devices use a film sensitive to x-rays to produce the images. Other technologies have been introduced to allow the image to be digitally acquired or to be projected on a screen and recorded on a media.

Image intensifiers: a specific component of an x-ray imaging system, which allows low intensity x-rays to be converted to a visible light output. The device contains a low absorbency/scatter input window, typically aluminum, input fluorescent screen, photocathode, electron optics, output fluorescent screen and output window. These parts are all mounted in a high vacuum environment within glass or more recently, metal/ceramic. It allows the viewer to more easily see the structure of the object being imaged than past fluorescent screens. This device was originally introduced in 1948. Viewing of the output was via mirrors and optical systems until the adaption of television systems in the 1960s. Additionally, the output was able to be captured on systems with a 100mm cut film camera using pulsed outputs from an x-ray tube similar to a normal radiographic exposure; Today CCD cameras are used.

Digital detectors: Digital detectors are an alternative to Image Intensifiers. The detector, roughly the same size of the image to be captured due to impracticability of focusing x-rays, converts x-rays into a charge than can be read and used to build the image. The most common technology is based on an indirect X-ray conversion process, using cesium iodide scintillators. It offers considerable advantages in radiography, angiography and fluoroscopy. The other method employs a direct converter such as selenium which is particularly suitable for mammography. Both flat detector technologies are based on amorphous silicon active pixel matrices.

Digital detectors facilitate the clinical workflow, ensure improved image quality which can help in reducing the dose at the expense of some energy as, to work correctly, they have to be kept at specific steady state temperature and therefore the x-ray device generally cannot be turned off. Digital detectors are also more fragile and are not suited for portable x-ray devices.

Battery powered x-ray equipment

Some x-ray equipment is powered by battery. The usage patterns and the energy consumption of such equipment are very different from the ones of mains powered x-rays. The SRI SC analysed only mains powered devices but discussions will continue in 2014 to include this specific category both in the measurement methodology and the ecodesign goal.
### 5.2. MARKET DATA

The COCIR companies\(^{51}\) in the X-ray sector represent a total turnover in Europe of \(\sqrt{740}\) \(^{52}\) million euros in 2016.

**Table 19: X-ray - EU\(^{53}\) market data**

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>X-Ray</td>
<td>(\sqrt{770}) M€</td>
<td>(\sqrt{843}) M€</td>
<td>(\sqrt{740}) M€</td>
</tr>
</tbody>
</table>

**Market coverage**

The X-ray modality comprises a lot of different categories. Unlike other imaging equipment in the scope of the SRI there is a big number of companies manufacturing x-ray devices. The actual SRI participating companies cover around 92% of the market for angiography systems, but less than 52% for all the remaining categories. Nonetheless COCIR believes that the best results can be achieved by developing concepts and methodologies that can encompass all x-ray categories. For this reason, the measurement methodology has been simplified to ensure it can be used to measure the energy consumption of any model, from huge angiography or interventional systems, to small mobile devices.

### 5.3. MEASUREMENT METHODOLOGY

The SRI Steering Committee started in 2013 to develop a methodology to measure the energy consumption of X-ray equipment, as there are no available standards that could be used.

The methodology ensures that:

- Results of the measurements are comparable and repeatable
- The measured energy consumption has a strict relation with the real energy consumption in everyday hospital use. Such information is important for hospitals and clinics to understand the real running cost of X-ray to plan and allocate correctly their budget
- The procedure does not involve disproportionate costs or resources
- The measurement results allow the determination of all the information that could be considered relevant for the SRI target setting process but could also be useful for X-ray users.

The methodology is available for download at COCIR website.

### 5.4. THE MEASUREMENT METHODOLOGY IN BRIEF

The SRI SC decided to use as functional unit a typical day of functioning of the X-ray scanner in hospital environment. According to expert judgment the time spent in scan mode is in the order of ‘seconds per day’ and therefore the energy consumption in scan mode is around 4% of the daily energy usage for the most energy intensive categories such as interventional or angiography x-ray.

The energy consumption in scan mode is also defined by many parameters and variables which should be defined for each specific x-ray category. The required complexity is not justified by the limited energy consumption during scans.

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\(^{51}\) The total turnover of COCIR Members in the x-ray sector is used here to avoid disclosing sensitive data each time a company joins the initiative, due to the large number of companies in this modality (only 6 in COCIR SRI).

\(^{52}\) COCIR Imaging Market Statistics source (SHARE).

\(^{53}\) Countries included: Estonia, Latvia, Lithuania, Bosnia, Bulgaria, Croatia, Czech Republic, Hungary, Poland, Romania, Serbia, Slovakia, Slovenia, Ukraine, Portugal, Spain, Denmark, Finland, Norway, Sweden, Ireland, UK, Austria, Belgium, France, Germany, Greece, Italy, Netherlands, Switzerland, Albania, Macedonia.
Therefore, the Steering Committee decided that the measurement methodology should not take into account scan mode, which would introduce a new layer of complexity to account for less than a 4% daily energy consumption.

**Functioning Modes**

Four modes have been defined for X-ray equipment.

**Off mode:** The system is shut down, AC mains off, according to the user manual. The system consumes no energy.

**Low-power:** The system functions into the minimum energy consumption state that the user can select according to the user manual.

**Ready-to-scan:** A state of the system when fully powered and ready to acquire the image.

**Scan mode:** A state of the system during scans. This mode includes tube rotor rotation, x-ray generation and generation of image.

The power consumption of X-ray in the four modes during a typical day is represented in figure 20.

![Interventional X-ray Daily Energy Consumption](image)

**Figure 20: Exemplary power consumption of MRI**

The daily energy consumption is calculated as the sum of the energy usage in the different modes. It has to be noted that certain technologies cannot be switched to off-mode (or it is highly recommended not to).
5.5. MODULARIZATION
The most important power consuming modules of the entire x-ray system have been identified.

| Tube and generator chain:      | X-ray tube and all the power supplies used to power the tube and all connected devices. |
| Detector/Image intensifier:    | Device reading the x-ray radiation and converting it into digital signal/image. |
| Power distribution unit and other power supplies | Subsystems required for electrical energy conversion and distribution |
| Computation, Controls:         | Units dedicated to the reconstruction and elaboration of the image, the control of the x-ray system. |
| Cooling:                      | A collection of subsystems required for thermal management of x-ray components, in particular the x-ray tube. |
| Patient table:                | Patient table with electric movement controls. |

5.6. ENERGY CONSUMPTION
X-ray devices are quite simple compared to the modalities analyzed so far by the SRI (MRI and CT) and the energy consumption is lower.
Considering the great variety of categories of x-rays, and the SRI Members market coverage, values are reported for complex x-rays such as angiography and interventional systems. For other categories lower power and energy consumption can be expected.

Table 20[^1]: Average x-ray energy consumption in different modes in case the x-ray is switched to LowPower mode overnight.

<table>
<thead>
<tr>
<th>Mode</th>
<th>Typical time in mode per day</th>
<th>Average power consumption</th>
<th>Average energy consumption per day</th>
</tr>
</thead>
<tbody>
<tr>
<td>LowPower</td>
<td>12 Hours</td>
<td>0,15 kW</td>
<td>1,8 kWh/d</td>
</tr>
<tr>
<td>Ready-to-scan</td>
<td>12 Hours</td>
<td>0,6 kW</td>
<td>7,3 kWh/d</td>
</tr>
<tr>
<td>Scan mode</td>
<td>Minutes</td>
<td>1,4 kW</td>
<td>-</td>
</tr>
</tbody>
</table>

The COCIR SRI focuses on angiography and interventional x-ray systems. The energy consumption for other x-ray devices is lower. For mammography systems, for instance, the power consumption in lowpower mode is lower than 0,1 kW.

5.7. IMPROVEMENT POTENTIAL
X-ray technology is not very different in principles from Computed Tomography. X-ray equipment are simpler than CT as the x-ray tube is not rotated by a gantry around the patient and the image is directly acquired without the need of complex engines for the reconstruction of a 3D image. CT scanners are also able to acquire hundreds of slices in one single examination.

[^1]: Values taken from the PE International Study: “Computed Tomography (CT) - Study on the potential for environmental improvement by the aspect of energy efficiency”, (2013) – Page 23
The COCIR expert group on x-ray analysed the x-ray technology and realized that the improvement potentials for modules are not different from the ones identified for CT equipment. Therefore the 26% improvement potential identified in LowPower mode can be reasonably assumed to be too high for X-ray keeping in mind that:

- X-ray equipment without digital detectors does not need to use energy for maintaining the steady state temperature, therefore the potential for improvement is lower.
- In 2014 it is expected only 15% of installed x-ray devices are equipped with digital detectors, and only high end stationary models. 45% of models sold in 2013/2014 are expected to be equipped with image intensifiers.
- The number of modules which are active is lower in x-ray, therefore there are lower chances to reduce the energy usage by switching of or to low power unnecessary modules.

As for CT, most x-ray devices are not switched off or to lowpower mode during night hours and therefore possible savings in such modes are not going to achieve any practical result.

5.8. USER BEHAVIOR

Considering the great variety of X-ray categories and different uses (from mobile c-arc to stationary interventional systems), it is impossible to identify a general user behaviour. Nonetheless considering the findings of previous projects it can be expected that most users do not profit of the low power or switch off functions of x-ray devices.

The same conclusion are contained in the study of the Danish Energy Saving Trust “Energy Efficiency in Hospitals and Laboratories“ where the energy consumption of x-ray and user behavior have been investigated:

"Energy savings can be achieved compared with current modes of operation. At present, in those hospitals involved in the project, X-ray equipment is always left in standby mode outside working hours because the medical staff considers that the reliability of the equipment will be impaired by frequent ON/OFF operation".

The following measurement table shows that over a 6 day measurement the x-ray device is never switched of maintaining a power absorption between 0,6 and 0,7 kW.

Figure 21: Power measurement of an x-ray device over a week in a Danish hospital
The study also concludes that:
"Secondly, despite the fact that most Danish hospitals have an impressive track record when it comes to implementing energy efficiency projects, the research showed that there is a significant, and additional energy saving potential. Many of the energy efficiencies are linked to working practice associated with the way in which medical equipment and laboratory equipment is operated."

5.9. INFLUENCING USER BEHAVIOUR
The SRI SC concluded that the greatest reduction in energy usage can be achieved by influencing the users’ behavior through proper education and information about the possible energy savings related to an environmental friendly use.

Already existing Off and LowPower modes could ensure an energy saving between 50,5% and 64,3% of daily energy consumption (see figure 22). Nonetheless such options are not widely used by users.

The use of Off mode during the night hours and weekend can save up to 3,45 MWh per year on average per equipment.

![Figure 22: Annual energy consumption of x-ray device per scenario](image-url)
APPENDIX VI

6. NUCLEAR IMAGING

6.1. GENERAL DESCRIPTION OF NUCLEAR IMAGING EQUIPMENT

In nuclear medicine imaging, the image is generated by using detectors (gamma cameras) which reads the gamma radiation emitted by a radio-isotope introduced in the body of the patient (orally or intravenously), whose properties bind it to certain types of tissues. This process is unlike a diagnostic X-ray, where external radiation is passed through the body to form an image.

Technologies

There are several techniques of nuclear imaging. This report focuses on positron emission tomography (PET) and single photon emission tomography (SPECT).

**SPECT** is very similar to conventional nuclear medicine planar imaging using a gamma camera but it is able to provide 3D information. Instead of just "taking a picture" of anatomical structures, a SPECT scan monitors level of biological activity at each place in the 3-D region analyzed. Emissions from the radionuclide indicate amounts of blood flow in the capillaries of the imaged regions.

To acquire SPECT images, the gamma camera is rotated around the patient. Projections are acquired at defined points during the rotation, typically every 3–6 degrees. In most cases, a full 360-degree rotation is used to obtain an optimal reconstruction. The time taken to obtain each projection is also variable, but 15–20 seconds is typical. This gives a total scan time of 15–20 minutes. Multi-headed gamma cameras can provide accelerated acquisition. For example, a dual-headed camera can be used with heads spaced 180 degrees apart, allowing two projections to be acquired simultaneously, with each head requiring 180 degrees of rotation. Triple-head cameras with 120-degree spacing are also used.

**PET** detects the gamma ray emission generated by a positron emitted by the decay of the radionuclide in the body of the patient. The positron travels in tissue for a short distance to a point where it interacts with an electron, annihilating each other producing a pair of annihilation (gamma) photons moving in opposite directions. These are detected when they reach a scintillator in the scanning device, creating a burst of light which is detected by photomultiplier tubes or silicon avalanche photodiodes (Si APD).

The technique depends on simultaneous or coincident detection of the pair of photons moving in opposite directions. Photons that do not arrive in temporal "pairs" (i.e. within a timing-window of a few nanoseconds) are ignored.
Combined Technologies
Nuclear imaging, combined with MRI and CT technologies are getting widespread for the additional benefits of adding precision of anatomic localization to functional imaging.

**PET/SPECT-CT** is a medical imaging technique combining a PET/SPECT scanner and a CT scanner, so that images acquired from both devices can be taken sequentially, in the same session, and combined into a single superposed (co-registered) image. Thus, functional imaging obtained by PET/SPECT, which depicts the spatial distribution of metabolic or biochemical activity in the body can be more precisely aligned or correlated with anatomic imaging obtained by CT scanning. Two- and three-dimensional image reconstruction may be rendered as a function of a common software and control system. The first PET-CT was installed in 1998.

**PET/SPECT-MRI** is a medical imaging technology that combines MRI soft tissue morphological imaging and PET/SPECT functional imaging. The technology combines the structural and functional characterization of tissue provided by MRI with the extreme sensitivity of PET/SPECT imaging of metabolism and tracking of uniquely labeled cell types or cell receptors. The first PET-MRI scanner was installed in 2010.

![Fig. 25: from left to right, two models of PET-CT and one model of PET-MRI](image)

6.2. MARKET DATA
The 355 companies participating in the SRI for the nuclear imaging sector represent a total turnover in Europe of 212 56 million euros in 2016, covering between 98% and 100% of the EU market.

In 2016 no PET scanners have been placed on the market as the technology has been replaced by PET-CT or PET-MRI.

In 2016, the market for SPECT scanners was around 60 mil €. While SPECT combined technologies are well established, there is still some market demand for SPECT scanners.

6.3. FUNCTIONING MODES

6.3.1. PET
In PET scanners, the PET gantry is always fully powered. Maintaining steady temperature of the scintillator crystals is fundamental as it would require otherwise a long time for the device to be ready to perform a scan. Unlike in MRI or Ct there is no emission of x-ray or RF during scan, therefore the energy usage is quite stable during the day.

**Off mode:** The system functions into the minimum energy consumption state that the user can select according to the user manual. As the PET gantry is expected to be running 24/7/365

---

55 GE Healthcare, Philips Healthcare, Siemens Healthcare
56 COCIR Imaging Market Statistics source (SHARE).
the system cannot be shut down to no-energy usage. Computers and accessories are turned off.

**Ready-to-scan/Scan mode**: All components are running. The power state/consumption is the same whether a scan is occurring or not.

### 6.3.2. SPECT
SPECT scanners can be turned off to zero energy consumption but they are normally left in low-power modes overnight as the detectors have to be maintained at a specific temperature. Warming up the scanner from Off mode takes a long time.

**Off**: The system is shut down, according to the user manual. The system consumes no energy.

**Low-power**: The system functions into the minimum energy consumption state that the user can select according to the user manual. No motion.

**Scan mode**: This mode represents the state of the system during scans with typical clinical motion and detectors acquiring data.

### 6.4. ENERGY CONSUMPTION
In both PET and SPECT systems, the energy consumption is quite stable as there is virtually no difference between scan mode and other modes. The main difference is the energy state of accessories such as computers and monitors which may be switched off during night time and week-ends.

### 6.5. PET: POWER USAGE AND DAILY ENERGY CONSUMPTION
Table 21 reports the power usage of the PET part of a PET-CT system and an average use scenario. The PET part absorbs around 4.3 kW in Off mode and a 16-20% in ready-to-scan/scan mode. This increase is due to the dedicated computers, up-to 5 computers with high output power supplies to handle multiple CPUs and GPUs. The daily energy consumption can be estimated in around 109 kWh/d.

Table 21: power usage of the PET part of a PET-CT scanner

<table>
<thead>
<tr>
<th>PET power usage</th>
<th>Off Mode</th>
<th>Ready-to-scan</th>
<th>Scan</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power Consumption (kW)</td>
<td>4.3</td>
<td>4.9</td>
<td>4.9</td>
</tr>
<tr>
<td>Daily usage (hours)</td>
<td>14</td>
<td>7</td>
<td>3</td>
</tr>
</tbody>
</table>

### 6.6. SPECT: POWER USAGE AND DAILY ENERGY CONSUMPTION
Table 22 reports the power usage of a SPECT scanner and an average use scenario. The SPECT absorbs 0.61 kW in Low-power mode and 0.63 in Scan-mode. The daily energy usage can be estimated around 15 kWh/d.

Table 22: power usage of a SPECT scanner not turned to off mode during the night

<table>
<thead>
<tr>
<th>SPECT : power usage</th>
<th>Off</th>
<th>Low-power</th>
<th>Scan</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power Consumption (kW)</td>
<td>0</td>
<td>0,61</td>
<td>0,63</td>
</tr>
<tr>
<td>Daily usage (hours)</td>
<td>0</td>
<td>16</td>
<td>8</td>
</tr>
</tbody>
</table>
**Fig. 26:** CT-PET power draw during a typical examination: IGEEOP Project (2009): “Indagine Sui Consumi Energetici Delle Grandi Apparecchiature Ospedaliere”
APPENDIX VII

7. METHODOLOGY FOR ASSESSING THE CIRCULARITY OF THE MEDICAL IMAGING DEVICES BUSINESS MODEL

In 2015, at the 4th Annual Forum on the COCIR SRI, COCIR presented its conclusions about reuse and refurbishment activities being one of the most significant contributions to reducing environmental impact of medical imaging devices. In 2015/2016 COCIR worked on a first methodology to estimate the circularity of the economic model. Such methodology aims at defining an index which can be later used for monitoring improvements. Additional work is needed to shape the methodology in such a way that the calculation of the index is based on data which can be easily obtained by companies with good approximation.

7.1. BENEFITS OF REFURBISHMENT

As already presented in many publications, refurbishment and repair activities brings significant benefits to the environment, patients and economy:

- Saves energy and CO2: by avoiding the production of new equipment, refurbishment contributes to save energy and CO2. DITTA\(^\text{57}\) estimates that around 30 MWh can be saved for each ton of refurbished medical devices.
- Prevents waste generation: DITTA estimates\(^\text{58}\) that in 2012 around 16.400 tons of used medical devices have been prevented from becoming waste, instead being shipped worldwide for refurbishment and repair.
- Saves resources and raw materials: medical devices make use of many scarce raw materials
- Contributes strongly to increased access to healthcare:
  - 20%-30% reduced cost for healthcare providers, while ensuring safety and high clinical performance.
  - Improvement of the age profile of installed equipment allowing hospitals with limited budget to substitute their old equipment.
  - Increase in quality of healthcare and safety for patients due to the reduction of the obsolescence of installed equipment.
- Contributes to safety of used medical devices which are restored to a point of safety and effectiveness comparable to when the device was new.
- Contributes to growth and economy. The refurbishment of medical equipment accounted for a global revenue of approximately 480 million euros in 2012\(^\text{59}\).

7.2. FROM LINEAR TO CIRCULAR: ROLE OF REFURBISHMENT

The European Union is aiming at moving the current “linear” model towards a circular one by means of modification of current legislation and introduction of new one. But moving from a linear model to a circular one is not a linear process. While it is clear that a perfect circular economy cannot be achieved, the degree of success depends on the implementation of all possible measures to prevent waste generation and use of resources from the design to the end of life phase, and to boost recycling of waste. The system complexity therefore increases non-linearly with the number of feedback loops which bring resources back to use diverting them from landfill, which is considered the end of the cycle.

\(^{57}\) DITTA: is the global voice for diagnostic imaging, radiation therapy, healthcare IT, electromedical and radiopharmaceutical manufacturers

\(^{58}\) Based on ad hoc DITTA internal survey on transboundary movements of used medical devices

\(^{59}\) DITTA Market Statistics
The best way to design a circular economy model for a specific industrial sector is to start from a linear one and to increase complexity of the model step by step adding loops which helps increasing resource efficiency level.

### 7.2.1. PERFECT LINEAR ECONOMY
A perfect linear economy\(^{60}\) is defined as a model where resources are used to make products which are discarded after the use phase without any recovery of materials.

![Perfect Linear Economy Diagram]

### 7.2.2. REAL LINEAR ECONOMY
In reality recycling of valuable fractions such as steel and copper has been in place for many years in almost any industrial sector. Nonetheless, in a linear economy, most of the waste generated is disposed of in landfill rather than being captured for recycling. This is represented by the red arrow leaving the “EOL” phase and ending directly in “LANDFILL”

![Real Linear Economy Diagram]

### 7.2.3. RECYCLING ECONOMY
With the introduction in EU of collective schemes for waste package (Waste Packaging Directive) and in 2005 of Extended Producer Responsibility schemes on electric and electronic products (WEEE Directive) and automotive (ELV Directive) the EU economy moved, for those sectors, towards a more recycling economy model. In this model most of the products are collected at the “EOL” phase and sent to recycling. Activities such as “preparation for reuse” can be in place but they will always be marginal for two reasons:
- Economy of the process
- Legislative barriers about reuse of recovered components in particular the chemical policy

Reuse may be an established activity but its role is anyway marginal. Reuse is driven by the economy of the process but reused products have to deal with consumer acceptance and the low cost of new products. Reuse activities are limited by legal barriers as the legislator has no attention to them.

---

\(^{60}\) PROD: production; EOL: end of life
The limits of this model are the recycling rate and the reuse of the recycled fractions:
- Technical limits to recycling rate
- Economic limits to recycling rate
- Economic value of recycled materials compared to virgin materials
- Purity/quality of recycled material
- Down-cycling rate

The down-cycling rate has to be seriously considered when defining the circularity of a specific industrial sector. When considering the economy in its entirety, down-cycling diverts resources from the landfill to other industrial sectors. Eventually such resources will have to be landfilled, thus leaving the cycle. In a specific industrial sector down-cycling has to be considered as the resources sent to recycling may not re-enter the cycle and new resources should be used despite the recycling rate.

The EC package on circular economy aims at increasing the way waste is managed, pushing towards a high level of collection and recycling. At the same time, the EC package also intends to promote real reuse of the recycled/recovered fractions by boosting the market for recovered materials. Those initiatives are more in line with the just examined “recycling economy model”.

### 7.2.4. SIMPLE CIRCULAR ECONOMY: INTRODUCING REUSE

The only way to move beyond the limits of a recycling economy is to prevent resources to become waste (a percentage is always destined to landfill). Reuse is a form of life time extension which allows resources to stay in the cycle longer before being sent to recycling. As long as resources are used, less new resources are needed to fill manufacturing and use needs. Simple reuse “as is”, without any additional processing, does not really extend the technical life time of a product. This practice simply prevents products to be discarded before they reach the end of their life, but also involves problems related with performances, reliability and safety. As such, simple reuse is going to be irrelevant.

Refurbishment is a specific form of reuse which restores performances, quality, safety and reliability to a state comparable to when the product/part was new. Refurbishment significantly extend the life time of products and parts solving the limits related to performances and safety.

In the model for the medical imaging sector “Preparation for reuse” is not considered as it is not a feasible option for medical devices. In fact, only parts or equipment which cannot be further refurbished and reused are discarded and sent to the “EOL” stage.
7.2.5. **THE MODEL**

This model includes three forms of reuse:

1. Reuse of used medical devices: used devices are refurbished and then sold.
2. Reuse of parts generated during refurbishment, repair or maintenance: such parts are refurbished and used for refurbishment, repair and maintenance.
3. Reuse of parts in manufacturing: reused parts are used in the manufacturing of new equipment.

![Diagram of the reusing cycle]

In the medical device sector experts estimate that up to 80% of certain modalities can be reused through refurbishment thus reducing significantly the quantity of material sent to “EOL” and then recycling.

**REUSE OF PARTS FOR MANUFACTURING NEW PRODUCTS**

Route 3, reusing parts in new products, could bring significant benefits in term of resource efficiency. This is indeed one of the best possible reuse activities. While reused and refurbished equipment are destined to a specific market with intrinsic limits, the reuse of parts into new equipment has unlimited potential.

The principle is already embedded in the EU legislation and international standards:

- **RoHS Directive**\(^{61}\): article 4.5 exempts recovered spare parts from RoHS when used to manufacture new equipment, under certain conditions.
- **ISO/IEC 24700:2004**: it specifies product characteristics for use in an original equipment manufacturer's declaration of conformity to demonstrate that a marketed product that contains reused components performs equivalent to new, meeting equivalent to new component specifications and performance criteria, and continues to meet all the safety and environmental criteria required by responsibly built products.
- **IEC IEC 62309:2004**: Introduces the concept to check the reliability and functionality of reused parts and their usage within new products. Also provides information and criteria about the tests/analysis required for products containing such reused parts, which are declared "qualified-as-good-as-new" relative to the designed life of the product. The purpose of this standard is to ensure by tests and analysis that the reliability and functionality of a new product containing reused parts is comparable to a product with only new parts.

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\(^{61}\) Directive 2011/65/EU of 8 June 2011 – L174/88
7.2.6. PERFECT CIRCULAR ECONOMY
In a perfect circular economy the use of natural resources is reduced to the minimum, just to supply the cycle with the resources which cannot be recovered by recycling due to physical or economic factors. The flowchart is the same but the input of resources is further reduced.

7.3. RESOURCE EFFICIENCY OF ECONOMY MODELS
The difference in the listed models, from the linear to the circular one, lies mostly in the amount of natural resources used per year. By defining the efficiency of the different operations such as recycling, refurbishment, reuse, etc and supposing the amount of products placed on the market equal to 100% it is possible to define a mass balance of the model and then to measure, at least in theory, the circularity of the model.

It is also important to assume that the installed base will not change over the year so a static system can be built. The variation in the installed base can be taken into account in a dynamic system but that would increase the complexity for no reason at this stage.

The COCIR SRI Steering Committee decided to define the Circularity Index as:

\[
C.I. = \frac{\text{Tons of reused equipment}}{\text{Tons of sold equipment}}
\]

Perfect circular economy - C.I. = 1
Perfect linear economy - C.I. = 0

In the following examples approximations based on expert judgment are used to quantify the different models for didactic reasons (refer to chapter 7.4 for additional information on how to calculate the index). The object is to show how reuse contributes to circularity to a great extent. No measurements have yet been taken by the SRI SC, therefore the final results may be quite different.

7.3.1. LINEAR ECONOMY
The models are now customized on the real model of the medical imaging sector. Resources are used to manufacture new equipment, parts and components which are used to maintain and repair the installed base. This distinction is important when dealing with a circular economy model.

\[
C.I. = \frac{16}{100} = 16\%
\]
7.3.2. REAL RECYCLING ECONOMY
In a real recycling economy most of the generated waste is collected and the recycling efficiency is as high as possible (technically and economically). The recycling efficiency also includes the rate of effective reuse of recycled material in the same loop (down-cycled resources are not considered).

C.I. = \frac{36}{100} = 36\%

7.3.3. PERFECT RECYCLING ECONOMY
In a perfect recycling economy almost 100% of the generated waste is collected and the recycling efficiency is as high as possible (technically and economically). The recycling efficiency also includes the rate of effective reuse of recycled material in the same loop (down-cycled resources are not considered).

C.I. = \frac{48}{100} = 48\%
7.3.4. **INTRODUCING REFURBISHMENT**

To move beyond the limits of recycling, reuse has to be implemented. In the medical imaging sector, the form reuse of whole equipment which ensures safety and performances is called refurbishment to differentiate from the simple reuse “as is”, or second hand market. Medical devices are bought back from owner by manufacturers or third parties to be refurbished and re-sold on the market.

![Diagram of refurbishment process]

C.I. = \( \frac{32 + 27}{100} = 59\% \)

7.3.5. **INTRODUCING REFURBISHMENT OF PARTS**

In the medical imaging sector, reuse of parts plays a fundamental role in ensuring:
- Servicing of the installed base
- Cost containment
- Prompt availability of parts for repair and servicing

Parts are sourced from different origins:
- Products under refurbishment
- Repair or maintenance activities

![Diagram of parts refurbishment process]

C.I. = \( \frac{35 + 27 + 5}{100} = 67\% \)
7.3.6. FULL SCALE CIRCULAR ECONOMY
A further and last step to improve the efficiency of the circular economy model is to increase the reuse rate of parts by using such recovered parts in the manufacturing on new equipment. This practice has been in place for a long time in the sector for specific high value/long life components, such as CT detectors, x-ray tubes or MRI magnets.

In a full scale circular economy model parts and equipment can be freely recovered, refurbished and reused whenever necessary without any unnecessary limitation provided by national, EU or international legislation.

\[
C.I. = \frac{32 + 27 + 5 + 10}{100} = 74\%
\]

7.4. MEASURING THE MEDICAL IMAGING SECTOR
In the full scale circular economy model the C.I. index is calculated as the sum of:

\[
C.I. = \frac{RM + PM + RE + RP}{POM} * 100
\]

Where:
RM: Mass of recycled materials
PM: Mass of parts re-used in manufacturing of new equipment
RE: Mass of refurbished equipment
RP: Mass of reused parts in repair and maintenance
POM: Mass of new products and parts placed on the market

Or
\[
C.I. = \frac{POM - LF}{POM} * 100
\]
Where:

LF: Mass sent to disposal

**RM: Mass of recycled materials**

The mass of recycled materials can be estimated knowing the total amount of waste sent to proper management and using an average value for the recycling rate. As companies in the medical imaging sector uses individual schemes to fulfil the obligations of the WEEE Directive, the recycling rate can be sourced from recyclers of medical devices. In reality the calculation could be more accurate but counting only the fractions which is really reused (metals) and excluding the fraction which is down-cycled (e.g. plastic)

\[ RM = (W \times \eta) \]

Where

\( \eta \) : recycling rate

**RP: Mass of reused parts in repair and maintenance**

While it may be possible to estimate the number of parts used in repair and maintenance of the installed base it is more difficult to know the mass as the weight of parts is not used and due to the heterogeneity, an average weight is meaningless. On the other end it may be assumed that for each recovered part used to repair an installed medical device, an equivalent part is discarded as waste or sent to refurbishment. For parts discarded as waste, the weight should be known. The complexity is represented by the numerous service providers who normally take care of installed equipment and which, sometimes are not related to the OEM. For parts shipped for refurbishment the weight of the shipments should be known for each shipment.

**PM: Total mass of reused parts in production of new equipment**

This data should be known to companies. In particular, after July 2014, this activity is mostly forbidden and therefore close to zero, with the exception of RoHS compliant spare parts:

- No RoHS substances or
- Applications covered by exemptions

**RE: Total mass of refurbished equipment placed on the market**

This data can be collected direct from companies. COCIR Market SHARE system already collects the sales volume in units. By defining an average weight an estimation of the mass can be possible.
APPENDIX VIII

8. EPD - ENVIRONMENTAL PRODUCT DECLARATION
The format developed by participating companies enables the individual member companies to systematically use the Environmental Product Declaration to communicate the most significant aspect(s) that the SRI sets for each targeted modality. Additional information on the EPD is available in the SRI Methodology Appendix 5.

<table>
<thead>
<tr>
<th>MINIMUM EPD REQUIREMENT including SRI targets and aspects</th>
</tr>
</thead>
<tbody>
<tr>
<td>SRI CONTENT - mandatory</td>
</tr>
<tr>
<td>SRI</td>
</tr>
<tr>
<td>Energy use according to specific scenarios and operating conditions</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Strongly recommended:</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Energy related</strong></td>
</tr>
<tr>
<td>CO₂ footprint in use phase according to specific scenarios and operating conditions</td>
</tr>
<tr>
<td>Environmentally relevant content/weight information</td>
</tr>
<tr>
<td>Product</td>
</tr>
<tr>
<td>Weight of product</td>
</tr>
<tr>
<td>Type and number of batteries</td>
</tr>
<tr>
<td>Relevant materials content - e.g. list RoHS, REACH SVHC, BoMcheck...</td>
</tr>
<tr>
<td>Packaging</td>
</tr>
<tr>
<td>Weight</td>
</tr>
<tr>
<td>composition</td>
</tr>
<tr>
<td>recyclable material content</td>
</tr>
<tr>
<td>Additional Ecologically relevant information</td>
</tr>
<tr>
<td>End of life aspects</td>
</tr>
<tr>
<td>refurbishing program available for the system</td>
</tr>
<tr>
<td>re-use of components program available for the system components</td>
</tr>
<tr>
<td>cleaning disinfection needed yes/no, if yes which chemicals</td>
</tr>
<tr>
<td>Information for user and recyclers (includes WEEE recycling passport info)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Optional</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Energy related</strong></td>
</tr>
<tr>
<td>Patient throughput for standard operation or energy per analysis</td>
</tr>
<tr>
<td>Waste during normal use (hazardous /non-hazardous/predefined categories?)</td>
</tr>
<tr>
<td>Emissions during normal use (hazardous /non-hazardous?)</td>
</tr>
<tr>
<td>Additional Ecologically relevant information</td>
</tr>
<tr>
<td>heat dissipation output - operating, stand-by, cooling,</td>
</tr>
<tr>
<td>start up time</td>
</tr>
<tr>
<td>Additional relevant information</td>
</tr>
<tr>
<td>Power and material saving options (e.g. to previous product)</td>
</tr>
</tbody>
</table>

⁶² This unit depends on the targeted environmental aspect. In this case it is “energy use” and thus “kWh”. It might also be a material, e.g. “copper” and thus “kg”.

### Additional End of Life aspects
- material recycling possibilities, Cradle to Cradle: describe
- ease of dismantling: describe

### Life cycle impact - company specific -
- % impact per LCA phase (e.g. 10% materials, 80% use; specify LCIA method): %/ describe

### Noise, radiation, vibration, EMC
- dB(A)

### Other Health and Safety related information
- voluntary actions (product and/or company specific): describe
- CSR related: describe
- Environmental education: describe
APPENDIX IX

9. INDEPENDENT PRACTITIONER’S LIMITED ASSURANCE REPORT

To the European Coordination Committee of the Radiological, Electromedical and Healthcare IT Industry ("COCIR"), the SRI Steering Committee Secretariat of the Self-Regulatory Initiative ("SRI") for Medical Imaging Equipment under the Ecodesign Directive, Brussels.

We have been engaged to perform a limited assurance engagement on selected data marked with ✓ in the Status Report of the SRI for Medical Imaging Equipment under the Ecodesign Directive (hereafter the "SRI Status Report"), Brussels, for the year 2016 in the context of the reporting requirements of the SRI.

Management’s Responsibility

The SRI Steering Committee Secretariat of the SRI is responsible for the preparation and presentation of the SRI Status Report in accordance with the six step methodology as stated in the SRI V1 (for ultrasound imaging equipment) and SRI V3 (for MRI, CT and future modalities) (hereafter the “SRI six step methodology”):

Step 1: Gather baseline data
Step 2: Prioritization and selection of next modality
Step 3: Identification of significant environmental aspect(s) for the selected modality
Step 4: Derive environmental targets and objectives for the selected modality
Step 5: Implementation into company processes
Step 6: Monitoring and reporting

and for the selection of the information to be assessed.

This responsibility includes the selection and application of appropriate methods to prepare the SRI Status Report as well as the use of assumptions and estimates for individual SRI disclosures which are reasonable in the circumstances. Furthermore, the responsibility includes designing, implementing and maintaining systems and processes relevant for the preparation of the SRI Status Report, which is free of material misstatements due to intentional or unintentional errors.

Audit Firm’s Independence and Quality Control

We have complied with the German professional provisions regarding independence as well as other ethical requirements.

The audit firm applies the national legal requirements and professional standards – in particular the Professional Code for German Public Auditors and German Chartered Auditors ("Berufssatzung für Wirtschaftsprüfer und vereidigte Buchprüfer": “BS WP/vBP”) as well as the Standard on Quality Control 1 published by the Institut der Wirtschaftsprüfer (Institute of Public Auditors in Germany; IDW): Requirements to quality control for audit firms (IDW Qualitätssicherungsstandard 1: Anforderungen an die Qualitätssicherung in der Wirtschaftsprüferpraxis - IDW QS 1) – and accordingly maintains a comprehensive system of quality control including documented policies and procedures regarding compliance with ethical requirements, professional standards and applicable legal and regulatory requirements.

Practitioner’s Responsibility

Our responsibility is to express a limited assurance conclusion on the data marked with ✓ in the SRI Status Report based on our work performed. Within the scope of our engagement we did not perform a substantive audit on external sources of information or expert opinions, used to prepare and referred to in the SRI Status Report.

We conducted our work in accordance with the International Standard on Assurance Engagements (ISAE) 3000 (Revised): “Assurance Engagements other than Audits or Reviews of Historical Financial Information” published by IAASB. This Standard requires that we plan and perform the assurance engagement to obtain limited assurance whether any matters have
come to our attention that cause us to believe that the data marked with ✓ in the SRI Status Report has not been prepared, in all material respects, in accordance with the SRI six step methodology.

In a limited assurance engagement the evidence-gathering procedures are more limited than for a reasonable assurance engagement and therefore significantly less assurance is obtained than in a reasonable assurance engagement. The procedures selected depend on the practitioner’s judgement. This includes the assessment of the risks of material misstatements of the data marked with ✓ in the SRI Status Report with regard to the above mentioned SRI six step methodology.

Within the scope of our work we performed amongst others the following procedures:

- Inquiries of personnel at the SRI Steering Committee Secretariat (COCIR) responsible for the preparation of the SRI Status Report regarding the process to prepare the report and the underlying internal control system;
- Inspection and sample testing of the systems and process documentation for collection, analysis and aggregation of the selected data;
- Recalculation of the aggregation and KPIs calculation for selected data;
- Analytical procedures on selected data;
- Inspection of documents regarding the description of the SRI six step methodology (SRI V3) and its application to MRI as detailed in ‘Magnetic resonance Equipment (MRI) - Study on the potential for environmental improvement by the aspect of energy efficiency’ and ‘Magnetic Resonance – Measurement of energy consumption’.

Conclusion

Based on our limited assurance engagement, nothing has come to our attention that causes us to believe that the data marked with ✓ in the SRI Status Report of the SRI for Medical Imaging Equipment under the Ecodesign Directive for the year 2016 has not been prepared, in all material respects, in accordance with the SRI six step methodology.

Emphasis of Matter – Recommendations

Without qualifying our conclusion above, we make the following recommendations for the further development of the reporting of the Self-Regulatory Initiative for Medical Imaging Equipment under the Ecodesign Directive:

- Further improvement of effectiveness and documentation of controls

Restriction on Use and Distribution

We issue this report on the basis of the engagement agreed with COCIR. The assurance engagement has been performed for purposes of the SRI for Medical Imaging Equipment under the Ecodesign Directive and is solely intended to inform SRI for Medical Imaging Equipment under the Ecodesign Directive about the results of the assurance engagement. The report is not intended for any third parties to base any (financial) decision thereon. Our responsibility lies only with COCIR. We do not assume any responsibility towards third parties.

Munich, 7 September 2017
PricewaterhouseCoopers GmbH
Wirtschaftsprüfungsgesellschaft

Hendrik Fink
Wirtschaftsprüfer
(German Public Auditor)
General Engagement Terms
for
Wirtschaftsprüfer und Wirtschaftsprüfungsgesellschaften
[German Public Auditors and Public Audit Firms]
as of January 1, 2017

1. Scope of application

(1) These engagement terms apply to contracts between German Public Auditors (Wirtschaftsprüfer) or German Public Audit Firms (Wirtschaftsprüfungsgesellschaften) – hereinafter collectively referred to as “German Public Auditors” – and their engaging parties for assurance services, tax advisory services, advice on business matters and other engagements except as otherwise agreed in writing or prescribed by a mandatory rule.

(2) Third parties may derive claims from contracts between German Public Auditors and engaging parties only when this is expressly agreed or results from mandatory rules. In relation to such claims, these engagement terms also apply to these third parties.

2. Scope and execution of the engagement

(1) Object of the engagement is the agreed service – not a particular economic result. The engagement will be performed in accordance with the German Principles of Proper Professional Conduct (Grundsätze ordnungsmäßiger Berufsausübung). The German Public Auditor does not assume any management functions in connection with his services. The German Public Auditor is not responsible for the use or implementation of the results of his services. The German Public Auditor is entitled to make use of competent persons to conduct the engagement.

(2) Except for assurance engagements (betriebswirtschaftliche Prüfungen), the consideration of foreign law requires an express written agreement.

(3) If circumstances or the legal situation change subsequent to the release of the final professional statement, the German Public Auditor is not obliged to refer the engaging party to changes or any consequences resulting therefrom.

3. The obligations of the engaging party to cooperate

(1) The engaging party shall ensure that all documents and further information necessary for the performance of the engagement are provided to the German Public Auditor on a timely basis, and that he is informed of all events and circumstances that may be of significance to the performance of the engagement. This also applies to those documents and further information, events and circumstances that first become known during the German Public Auditor’s work. The engaging party will also designate suitable persons to provide information.

(2) Upon the request of the German Public Auditor, the engaging party shall confirm the completeness of the documents and further information provided as well as the explanations and statements, in a written statement drafted by the German Public Auditor.

4. Ensuring independence

(1) The engaging party shall refrain from anything that endangers the independence of the German Public Auditor’s staff. This applies throughout the term of the engagement, and in particular to offers of employment or to assume an executive or non-executive role, and to offers to accept engagements on their own behalf.

(2) Were the performance of the engagement to impair the independence of the German Public Auditor of related firms, firms within its network, or such firms associated with him, to which the independence requirements apply in the same way as to the German Public Auditor in other engagement relationships, the German Public Auditor is entitled to terminate the engagement for good cause.

5. Reporting and oral information

To the extent that the German Public Auditor is required to present results in writing as part of the work in executing the engagement, only that written work is authoritative. Drafts are non-binding. Except as otherwise agreed, oral statements and explanations by the German Public Auditor are binding only when they are confirmed in writing. Statements and information of the German Public Auditor outside of the engagement are always non-binding.

6. Distribution of a German Public Auditor’s professional statement

(1) The distribution to a third party of professional statements of the German Public Auditor (results of work or extracts of the results of work whether in draft or in a final version) or information about the German Public Auditor acting for the engaging party requires the German Public Auditor’s written consent, unless the engaging party is obligated to distribute or inform due to law or a regulatory requirement.

(2) The use by the engaging party for promotional purposes of the German Public Auditor’s professional statements and of information about the German Public Auditor acting for the engaging party is prohibited.

7. Deficiency rectification

(1) In case there are any deficiencies, the engaging party is entitled to specific subsequent performance by the German Public Auditor. The engaging party may reduce the fees or cancel the contract for failure of such subsequent performance, for subsequent non-performance or unjustified refusal to perform subsequently, or for unconscionability or impossibility of subsequent performance. If the engagement was not commissioned by a consumer, the engaging party may only cancel the contract due to a deficiency if the service rendered is not relevant to him due to failure of subsequent performance, to subsequent non-performance, to unconscionability or impossibility of subsequent performance. No. 9 applies to the extent that further claims for damages exist.

(2) The engaging party must assert a claim for the rectification of deficiencies in writing (Textform) [Translators Note: The German term “Textform” means in written form, but without requiring a signature] without delay. Claims pursuant to paragraph 1 not arising from an intentional act expire after one year subsequent to the commencement of the time limit under the statute of limitations.

(3) Apparent deficiencies, such as clerical errors, arithmetical errors and deficiencies associated with technicalities contained in a German Public Auditor’s professional statement (long-form reports, expert opinions etc.) may be corrected – also versus third parties – by the German Public Auditor at any time. Misstatements which may call into question the results contained in a German Public Auditor’s professional statement entitle the German Public Auditor to withdraw such statement – also versus third parties. In such cases the German Public Auditor should first hear the engaging party, if practicable.

8. Confidentiality towards third parties, and data protection

(1) Pursuant to the law (§ [Article] 323 Abs 1 [paragraph 1] HGB [German Commercial Code: Handelsgesetzbuch]), § 43 WPO [German Law regulating the Profession of Wirtschaftsprüfer: Wirtschaftsprüferordnung], § 203 StGB [German Criminal Code: Strafgesetzbuch]) the German Public Auditor is obligated to maintain confidentiality regarding facts and circumstances related to his professional work, unless the engaging party releases him from this confidentiality obligation.

(2) When processing personal data, the German Public Auditor will observe national and European legal provisions on data protection.

9. Liability

(1) For legally required services by German Public Auditors, in particular audits, the respective legal limitations of liability, in particular the limitation of liability pursuant to § 323 Abs. 2 HGB, apply.

(2) Insofar neither a statutory limitation of liability is applicable, nor an individual contractual limitation of liability exists, the liability of the German Public Auditor for claims for damages of any other kind, except for damages resulting from injury to life, body or health as well as for damages that constitute a duty of replacement by a producer pursuant to § 1 ProdHaftG [German Product Liability Act: Produkthaftungsgesetz], for an individual case of damages caused by negligence is limited to € 4 million pursuant to § 54 a Nr. 2 WPO.

(3) The German Public Auditor is entitled to invoke demurs and defenses based on the contractual relationship with the engaging party also towards third parties.
(4) When multiple claimants assert a claim for damages arising from an existing contractual relationship with the German Public Auditor due to the German Public Auditor’s negligent breach of duty, the maximum amount stipulated in paragraph 2 applies to the respective claims of all claimants collectively.

(5) An individual case of damages within the meaning of paragraph 2 also exists in relation to a uniform damage arising from a number of breaches of duty. The individual case of damages encompasses all consequences from a breach of duty regardless of whether the damages occurred in one year or in a number of successive years. In this case, multiple acts or omissions based on the same source of error or on a source of error of an equivalent nature are deemed to be a single breach of duty if the matters in question are legally or economically connected to one another. In this event the claim against the German Public Auditor is limited to € 5 million. The limitation to the fivefold of the minimum amount insured does not apply to compulsory audits required by law.

(6) A claim for damages expires if a suit is not filed within six months subsequent to the written refusal of acceptance of the indemnity and the engaging party has been informed of this consequence. This does not apply to claims for damages resulting from scienter, a culpable injury to life, body or health as well as for damages that constitute a liability for replacement by a producer pursuant to § 1 ProdHaRg. The right to invoke a plea of the statute of limitations remains unaffected.

10. Supplementary provisions for audit engagements

(1) If the engaging party subsequently amends the financial statements or management report audited by a German Public Auditor and accompanied by an auditor’s report, he may no longer use this auditor’s report.

If the German Public Auditor has not issued an auditor’s report, a reference to the audit conducted by the German Public Auditor in the management report or any other public reference is permitted only with the German Public Auditor’s written consent and with a wording authorized by him.

(2) If the German Public Auditor revokes the auditor’s report, it may no longer be used. If the engaging party has already made use of the auditor’s report, then upon the request of the German Public Auditor he must give notification of the revocation.

(3) The engaging party has a right to five official copies of the report. Additional official copies will be charged separately.

11. Supplementary provisions for assistance in tax matters

(1) When advising on an individual tax issue as well as when providing ongoing tax advice, the German Public Auditor is entitled to use as a correct and complete basis the facts provided by the engaging party – especially numerical disclosures; this also applies to bookkeeping engagements. Nevertheless, he is obligated to indicate to the engaging party any errors he has identified.

(2) The tax advisory engagement does not encompass procedures required to observe deadlines, unless the German Public Auditor has explicitly accepted a corresponding engagement. In this case the engaging party must provide the German Public Auditor with all documents required to observe deadlines – in particular tax assessments – on such a timely basis that the German Public Auditor has an appropriate lead time.

(3) Except as agreed otherwise in writing, ongoing tax advice encompasses the following work during the contract period:

a) preparation of annual tax returns for income tax, corporate tax and business tax, as well as wealth tax returns, namely on the basis of the annual financial statements, and on other schedules and evidence documents required for the taxation, to be provided by the engaging party
b) examination of tax assessments in relation to the taxes referred to in (a)
c) negotiations with tax authorities in connection with the returns and assessments mentioned in (a) and (b)
d) support in tax audits and evaluation of the results of tax audits with respect to the taxes referred to in (a)
e) participation in petition or protest and appeal procedures with respect to the taxes mentioned in (a).

In the aforementioned tasks the German Public Auditor takes into account material published legal decisions and administrative interpretations.

(4) If the German Public auditor receives a fixed fee for ongoing tax advice, the work mentioned under paragraph 3 (d) and (e) is to be remunerated separately, except as agreed otherwise in writing.

(5) Insofar the German Public Auditor is also a German Tax Advisor and the German Tax Advice Remuneration Regulation (Steuerberatungsvergütungsverordnung) is to be applied to calculate the remuneration, a greater or lesser remuneration than the legal default remuneration can be agreed in writing (Textform).

(6) Work relating to special individual issues for income tax, corporate tax, business tax, valuation assessments for property units, wealth tax, as well as all issues in relation to sales tax, payroll tax, other taxes and duties requires a separate engagement. This also applies to:

a) work on non-recurring tax matters, e.g. in the field of estate tax, capital transactions tax, and real estate sales tax;
b) support and representation in proceedings before tax and administrative courts and in criminal tax matters;
c) advisory work and work related to expert opinions in connection with changes in legal form and other re-organizations, capital increases and reductions, insolvency related business reorganizations, admission and retirement of owners, sale of a business, liquidations and the like, and
d) support in complying with disclosure and documentation obligations.

(7) To the extent that the preparation of the annual sales tax return is undertaken as additional work, this includes neither the review of any special accounting prerequisites nor the issue as to whether all potential sales tax allowances have been identified. No guarantee is given for the complete compilation of documents to claim the input tax credit.

12. Electronic communication

Communication between the German Public Auditor and the engaging party may be via e-mail. In the event that the engaging party does not wish to communicate via e-mail or sets special security requirements, such as the encryption of e-mails, the engaging party will inform the German Public Auditor in writing (Textform) accordingly.

13. Remuneration

(1) In addition to his claims for fees, the German Public Auditor is entitled to claim reimbursement of his expenses; sales tax will be billed additionally. He may claim appropriate advances on remuneration and reimbursement of expenses and may make the delivery of his services dependent upon the complete satisfaction of his claims. Multiple engaging parties are jointly and severally liable.

(2) If the engaging party is not a consumer, then a set-off against the German Public Auditor’s claims for remuneration and reimbursement of expenses is admissible only for undisputed claims or claims determined to be legally binding.

14. Dispute Settlement

The German Public Auditor is not prepared to participate in dispute settlement procedures before a consumer arbitration board (Verbraucherschlichtungsstelle) within the meaning of § 2 of the German Act on Consumer Dispute Settlements (Verbraucherstreitbeilegungsgesetz).

15. Applicable law

The contract, the performance of the services and all claims resulting therefrom are exclusively governed by German law.