EXPLANATORY MEMORANDUM TO

THE CONTROL OF ARTIFICIAL OPTICAL RADIATION AT WORK
REGULATIONS 2010

2010 No. 1140

1. This explanatory memorandum has been prepared by The Health and Safety Executive (HSE) on behalf of the Department for Work and Pensions and is laid before Parliament by Command of Her Majesty.

This memorandum contains information for the Joint Committee on Statutory Instruments.

2. Purpose of the instrument

2.1 These Regulations place duties on employers to protect workers from the risks from hazardous sources of artificial light (artificial optical radiation) in the workplace.


3. Matters of special interest to the Joint Committee on Statutory Instruments

3.1 The Regulations rely on the power in paragraph 1A of Schedule 2 to the European Communities Act 1972 so that the Regulations can refer to Annexes of the Directive "as amended from time to time". This is because Article 10 of the Directive allows technical amendments to be made to the Annexes so they may be updated in the future.

3.2 The use of this enabling power is considered by the Secretary of State to be appropriate and expedient. Any amendments to the Annexes will be technical in nature, and the use of the power avoids the need to bring forward new legislation that would simply reproduce or refer to the amended Annexes. In this way, references in the Regulations to the Annexes of the Directive will always reflect the most up to date version of the Annexes without the need for time consuming amendments that will simply be technical in nature.

3.3 HSE will publicise any proposals to amend the Annexes on its website and will amend its guidance in advance of any technical changes being made.

4. Legislative Context

4.1 The Directive is the nineteenth daughter directive of the health and safety Framework Directive 89/391/EEC, and the fourth and last in a sequence of so-called “physical agents” Directives. The physical agents Directives are derived from an amendment of the European Commission’s original proposal in 1993 for a single

- **Scrutiny history**

  4.2 Under the Irish Presidency, the EU developed a proposal for the Directive that was presented to the EU Council in July 2004 and negotiated with the Council’s Social Questions Working Group.

  4.3 The proposal for the Directive and the accompanying explanatory memorandum were considered by the House of Commons and House of Lords’ Select Committees on European Scrutiny respectively in November 2004. The proposal was held under scrutiny by the Commons pending sight of the regulatory impact assessment (“RIA”) and referred by the Lords to their Sub-Committee G.

  4.4 The Minister of State for Work, Jane Kennedy, forwarded the RIA by letter to the Lords and Commons scrutiny committees on 15 November 2004. The House of Commons considered the letter and the RIA in December 2004 and referred the RIA for debate in the House of Commons European Standing Committee B.

  4.5 Whilst the House of Lords Sub-Committee G considered the EM and RIA on 17 November 2004 and released it from scrutiny, the Committee chairman, Lord Grenfell, wrote to the Minister on 24 November 2004 and 7 April 2005 expressing concerns about the requirement in the proposed Directive to assess risks from exposure to natural sources (essentially sunlight); the impact costs and the fact that it had not been possible to consult small organisations because of the pace at which the EU negotiations were being conducted. All these matters were subsequently addressed, following the amendment to the proposed Directive by the European Parliament at its second reading, as referred to below.

  4.6 The House of Commons European Standing Committee B debated the proposal on 24 January 2005 and voted to support the Government’s position on the proposal.

  4.7 The proposed Directive was passed to the European Parliament for its second reading in September 2005. This resulted in amendments to remove the sunlight provisions.

- **Transposition**

  4.8 A Transposition Note is attached at Appendix 1 of this memorandum.

  4.9 The Directive has been implemented in the most proportionate manner which meets the objectives of the Directive while avoiding unnecessary burdens on employers. Employers are only obliged to carry out a detailed risk assessment (which includes assessing, and if necessary measuring, levels of exposure to artificial light) under the Regulations where an employer carries out work which could expose its employees to levels of artificial optical radiation that could create a reasonably foreseeable risk of adverse health effects to the eyes or skin and where those risks have not already been eliminated or controlled.
4.10 The Regulations also fulfil the obligation under Article 9 of the Directive to provide for adequate penalties for infringement of the Regulations that are effective, proportionate and act as a deterrent. Section 33 of Health and Safety at Work Act 1974 will apply and this makes it an offence for employers to contravene any health and safety regulations.

5. Territorial Extent and Application

This instrument applies to Great Britain. Separate regulations are to be made to implement the Directive in Northern Ireland and Gibraltar.


As the instrument is subject to negative resolution procedure and does not amend primary legislation, no statement is required.

7. Policy background

- What is being done and why

7.1 The aim of the Directive is to protect workers from the risks arising from exposure to sources of artificial light that the Directive describes as artificial optical radiation. Artificial optical radiation can occur as ultraviolet light, infrared radiation and as laser beams with the risks from exposure dependent on the type and intensity of the light and the parts of the body exposed. The skin and the eyes are the areas of the body most at risk.

7.2 The Regulations require employers to consider whether there are any foreseeable risks of adverse health effects to the eyes or skin of their workers as a result of exposure to artificial light. HSE guidance will help employers to do this. Depending on the outcome of that assessment, and if they haven’t taken steps to eliminate, or reduce to as low as is reasonably practicable these risks (HSE guidance will help them determine this), they will need to carry out a detailed risk assessment. Additional measures may also be necessary, such as developing an action plan, which will depend on the results of the risk assessment.

- Scale of the problem

7.3 The risk from exposure to intense forms of artificial light is not a significant problem within the UK. Many businesses will only have innocuous light sources such as those found in office environments. The number of businesses using intense sources of light that can be hazardous is estimated to be 80,000. These include research and education institutes using powerful lasers, metal fabrication work using welding equipment, printing processes using ultraviolet light and industries using furnaces. However, even within these sectors, there are few if any reports of ill health or injury. In the last 15 years there are estimated to have been less than 10 injuries that required workers to take more than 3 days off work reported to HSE. There have been no cases of work-related cataracts or neoplasia (new or abnormal tissue growth) attributed to exposure to artificial light reported by general practitioners or
occupational physicians. There have been 19 cases of work related neoplasia reported by consultant dermatologists through the Occupational Skin Surveillance Scheme that they attributed to exposure to artificial light. In 90% of these cases, the workers were involved in welding which generates high levels of ultraviolet light. However, it was not clear how many of these workers also worked outside or spent their leisure time outside – which would also increase their exposure to natural optical radiation (sunlight). There have also been 65 cases of heat cataracts (a prescribed industrial disease) compensated under the Industrial Injuries scheme since 1992 although no cases have been reported after 2002.

- **UK position on the Directive**

7.4 The Government did not oppose the Directive as it aims to standardise protection for workers across Europe and the Government wants to ensure that all workers benefit from this protection.

7.5 The initial draft Directive included additional provisions for assessing the risks of exposure from natural sources (essentially sunlight). However, the provisions on sunlight were subsequently removed by an amendment from the European Parliament. This move was endorsed by the UK because it was incompatible with the policy on better regulation.

- **Level of public interest**

7.6 There has only been very modest interest from the media in this Directive although some stakeholders have considered it to be an unnecessary piece of European legislation. Representatives from the manufacturing sector called for a review of the Directive and wrote to the Government and the Chair of the HSE prior to the launch of the consultation. However, HSE has continued to engage with these stakeholders to ensure that their concerns are addressed, in particular to avoid imposing unnecessary burdens where there is no risk of harm to workers.

7.7 Some stakeholders are already addressing the possible challenges posed by these new Regulations by identifying where there are gaps in exposure data and taking measures to plug them and engaging with key groups such as those associated with the entertainment sector to disseminate knowledge.

- **The need for legislation**

7.8 In transposing the Directive, the challenge was to devise a set of Regulations that would meet the Government’s obligations under the Directive without imposing unnecessary burdens on businesses where there is no risk of harm to workers. The Government also wanted to ensure that those businesses not already managing the risks effectively do more to protect their workers. The Regulations have been developed to meet these aims.

7.9 In addressing the challenge, three options were considered:

- Do nothing – continue to rely on existing regulatory provisions already in place in Great Britain
• Introduce a full set of new regulatory provisions to reproduce the full requirements of the Directive disregarding protections under existing regulatory provisions.
• Rely on protections under existing regulatory provisions where appropriate and introduce new regulatory provisions limited to new, specific requirements set by the Directive where necessary

7.10 The first option would have zero costs and benefits but would not meet the Government’s policy on proper transposition of the Directive and therefore was not viable.

7.11 The second option would have disregarded existing protections and would have placed unnecessary additional burdens on businesses without reducing the risks. This was not considered desirable.

7.12 The final option meets the objectives of the Directive – to protect workers from harm as a result of exposure from artificial light and is consistent with better regulation practice. In taking this approach, the Regulations have regard to existing protections and will essentially only have an impact on those employers who are dealing with hazardous sources of artificial light and are not already managing risks. This is reflected in the comparatively low costs associated with this Directive (see impact assessment below).

• Consolidation

7.13 As these Regulations do not amend any other legislation, the question of consolidation does not arise.

8. Consultation outcome

8.1 The consultation period started on 9 November 2009 and lasted for thirteen weeks, closing on 5 February 2010. The consultation package included the draft Regulations, an impact assessment, an initial first draft of guidance for employers and a questionnaire. It was sent to almost two hundred stakeholders including professional bodies, trade associations, medical and technical groups and employer and employee representatives. HSE received 60 responses with comments covering the Regulations, guidance and impact assessment.

8.2 There were twenty seven responses on the Regulations ranging from points of clarification to substantive comments. The draft Regulations that were issued for consultation referred back to obligations under the Management of Health and Safety at Work Regulations 1999 (the Management Regulations) and in particular the duty to risk assess under the Management Regulations. Several stakeholders reported that they were confused by this approach. Other comments were focussed on parts of the Directive that consultees did not consider to have been transposed in the Regulations. These included matters such as the appointment of competent persons, the confidentiality of records for people undergoing health surveillance and the exclusion of the eyes from the health surveillance requirements.
8.3 The Regulations have been re-drafted to improve clarity and understanding. Matters such as competent persons are already addressed within the Management Regulations, and following extensive in-house consultation on health surveillance for eyes, HSE has concluded that that there is no justification for its inclusion on the basis that there is currently no accepted eye examination that would provide meaningful information through routine monitoring. The regulations will require a worker whose eyes have been over-exposed to be offered a medical examination. The confidentiality of records is covered by the Data Protection Act 1998.

8.4 In relation to the guidance, almost all those who responded have called for more and better information, the inclusion of information on a wider range of sources, more information on light sources used in the entertainment industry and greater clarity. HSE fully accepts that much more needs to be done to produce guidance that is fit for purpose. HSE has met with a key official from the Professional Lighting and Sound Association which is the lead professional body for those who in particular supply lighting technology to the entertainment and communication sectors. Further discussions will take place on ways to disseminate knowledge about risk management to those who work in these industries. The output from these discussions will feed into HSE’s action plan to address the views of stakeholders and meet other relevant sector specific needs.

8.5 A substantive number of stakeholders also felt that the impact costs had been underestimated particularly in relation to the time required for familiarisation and information and training. HSE undertook a further review of the impact costs and the revised estimates are shown below.

9. **Guidance**

The Regulations will be supported by guidance from two sources. Firstly, HSE will develop and refine its initial guidance to assist employers in identifying what sources of light can cause harm and to outline what they should be doing to manage the risks. This will then be supported by additional sector specific guidance developed with and through others and will also be amended to alert stakeholders in advance of any technical changes to the Directive. Secondly extensive guidance, soon to be published although a draft is publicly available, has been developed at EU level for use by all Member States.

10. **Impact**

10.1 The impact on business, charities or voluntary bodies in relation to the first year costs is **£4.64 million** (best estimate with a range of £2.96m - £6.67m) and **£12.55 million** (best estimate with a range of £7.4m - £19.06m) at 10 year present value. These costs are spread fairly evenly between the implementation costs arising from screening, familiarisation and worker information to the policy costs associated with risk assessment and necessary action to reduce exposure.

10.2 These Regulations apply only to people at work but exclude the self-employed.
10.3 A summary of the Impact Assessment is shown below and the full impact assessment is attached at Appendix 2

<table>
<thead>
<tr>
<th>Implementation costs</th>
<th>First year costs</th>
<th>Ten year present value</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>£ million</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>Best estimate</td>
</tr>
<tr>
<td>Self screening</td>
<td>0.63</td>
<td>0.70</td>
</tr>
<tr>
<td>Familiarisation</td>
<td>0.47</td>
<td>0.70</td>
</tr>
<tr>
<td>Worker information</td>
<td>0.35</td>
<td>0.63</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Policy Costs</th>
<th>First year costs</th>
<th>Ten year present value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>£ million</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>Best estimate</td>
</tr>
<tr>
<td>Refresh risk assessment</td>
<td>0.25</td>
<td>0.42</td>
</tr>
<tr>
<td>New risk assessment (one off)</td>
<td>0.72</td>
<td>1.25</td>
</tr>
<tr>
<td>New risk assessment (recurring)</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Action to reduce exposure (time)</td>
<td>0.06</td>
<td>0.11</td>
</tr>
<tr>
<td>Action to reduce exposure (equipment)</td>
<td>0.39</td>
<td>0.72</td>
</tr>
<tr>
<td>Health surveillance</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>HSE costs</td>
<td>First year costs</td>
<td>Ten year present value</td>
</tr>
<tr>
<td></td>
<td>£ million</td>
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</tr>
<tr>
<td></td>
<td>Min</td>
<td>Best estimate</td>
</tr>
<tr>
<td>Produce guidance</td>
<td>0.09</td>
<td>0.10</td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td><strong>2.96</strong></td>
<td><strong>4.64</strong></td>
</tr>
</tbody>
</table>

11. **Regulating small business**

11.1 The legislation applies to small business.

11.2 To minimise the impact of the requirements on firms employing up to 20 people, the approach taken is two fold:

11.3 Firstly, employers do not need to carry out a detailed risk assessment if they are satisfied that the nature of their work does not expose any employees to risks of adverse health effects to the eyes or skin: either because they only have safe sources or hazardous sources that are well managed. In effect, this means that their responsibilities will be discharged by compliance with their existing duties.

11.4 Secondly, if the nature of their work is such that some further action is needed to meet the requirements of the new Regulations, then HSE’s guidance will help them.

12. **Monitoring & review**

12.1 HSE will liaise with key stakeholders and other professional bodies after the Regulations have come into force to assess whether they are meeting their desired purpose
and if the guidance is satisfactory in the light of experience. This will take place within two years of implementation by means of both formal and informal contact with dutyholders and representatives of professional bodies and trade associations.

12.2 The system for reporting accidents and injuries to HSE will also be reviewed to ensure that the existing extremely low incidence of reports of harm is maintained.

13. **Contact**

For more information on the statutory instrument contact Matthew Penrose at the Health and Safety Executive.

Tel: 0151 951 4909

Email: [matthew.penrose@hse.gsi.gov.uk](mailto:matthew.penrose@hse.gsi.gov.uk)
**TRANSPOSITION NOTE**


The Control of Artificial Optical Radiation at Work Regulations 2010 ("the Regulations") implement the Directive.

These Regulations do what is necessary to implement the Directive. The main elements of the Directive implemented in these Regulations are as follows:

<table>
<thead>
<tr>
<th>Article</th>
<th>Purpose</th>
<th>Implementation</th>
<th>Responsibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Sets out the aim and scope of the Directive.</td>
<td>Implicit in the Regulations as a whole.</td>
<td>Secretary of State</td>
</tr>
<tr>
<td>2</td>
<td>Definitions.</td>
<td>Regulation 1 and 3(8).</td>
<td>Secretary of State</td>
</tr>
<tr>
<td>3</td>
<td>Establishes the exposure limit values for non-coherent and laser radiation.</td>
<td>Regulation 1</td>
<td>Secretary of State</td>
</tr>
<tr>
<td>4</td>
<td>Requires employers to conduct a risk assessment to assess/measure/calculate levels of exposure to artificial optical radiation (AOR) in accordance with the prescribed methodology and with reference to particular criteria.</td>
<td>Regulation 3</td>
<td>Secretary of State</td>
</tr>
<tr>
<td>5</td>
<td>Requires employers to eliminate or reduce to a minimum risks of exposure to AOR and includes action and measures to take if the exposure limit values are exceeded.</td>
<td>Regulation 4</td>
<td>Secretary of State</td>
</tr>
<tr>
<td>6</td>
<td>Requires employers to ensure that workers receive any necessary information and training in relation to the outcome of the risk assessment, with particular reference to the matters listed.</td>
<td>Regulation 5</td>
<td>Secretary of State</td>
</tr>
<tr>
<td>7</td>
<td>Consultation and participation of workers in accordance</td>
<td>The Safety Representatives &amp; Secretary of State</td>
<td></td>
</tr>
</tbody>
</table>

Safety Committees Regulations 1977 & Health and Safety (Consultation with Employees) Regulations 1996 already provide for consultation with, and the participation of, employees in matters relating to health and safety.

8.1 Member States to adopt measures to ensure appropriate health surveillance.

Regulation 6(1) The Secretary of State

8.2-8.3 Prescribes arrangements for health surveillance and access to health surveillance documents.

Regulation 6(2)-(5) The Secretary of State

8.4 Establishes obligation on employer to make available a medical examination where exposure above the limit values is detected and as a result of any adverse health effect of exposure found following health surveillance. In both cases there is an obligation on employers to review the risk assessment and measures taken in accordance with Article 5 and to carry out continued health surveillance and if necessary a medical examination in accordance with the prescribed arrangements.

Regulations 6(6) and (7) The Secretary of State

9 Establishes obligation for Member States to establish a penalty regime for breaches of the Directive.

The Health and Safety at Work etc. Act 1974 provides for penalties for breach of health and safety regulations.

The Secretary of State

10 and 11 Prescribe the legislative procedure for amendments to the annexes of the Directive.

Transposition not required.


12 Prescribes obligations for Member States to provide a report to the European Commission on the practical

Transposition not required. In any event this requirement was repealed with effect
implementation of the Directive and for the Commission to inform the European Parliament, the Council and the prescribed committees of the content of these reports and other associated information.

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<tr>
<td>13</td>
<td>European Commission to produce a practical guide.</td>
<td>Transposition not required.</td>
</tr>
<tr>
<td>14</td>
<td>Prescribes measures for Member States to implement the Directive.</td>
<td>Transposition not required.</td>
</tr>
<tr>
<td>15</td>
<td>Provides for the entry into force of the Directive</td>
<td>Transposition not required.</td>
</tr>
<tr>
<td>Annex I</td>
<td>Prescribes the exposure limit values for non-coherent radiation.</td>
<td>Regulation 1.</td>
</tr>
<tr>
<td>Annex II</td>
<td>Prescribes the exposure limit values for laser radiation.</td>
<td>Regulation 1</td>
</tr>
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</table>

from 27.6.07 by 2007/30/EC
What is the problem under consideration? Why is government intervention necessary?

Businesses have a duty to ensure that hazardous sources of Artificial Optical Radiation (AOR) in their workplace (e.g. lasers and UV light) are managed so that the health and safety of their workers is protected. The Control of Artificial Optical Radiation at Work Regulations 2010 will ensure that those businesses not already doing this take action to ensure the risks to their workers are reduced to as low a level as is reasonably practicable; those businesses where workers are already at low risk will not need to do anything more. The Regulations will transpose the specific requirements of the European Physical Agents (Artificial Optical Radiation) Directive.

What are the policy objectives and the intended effects?

To transpose a Directive from the European Union which prescribes control measures to protect the health and safety of workers across European Member States from hazardous sources of AOR. The policy objectives are to (a) ensure that all workers in Great Britain are sufficiently protected (b) to meet the Government's Treaty obligations to transpose the Directive and (c) to meet these in a proportionate way which minimises unnecessary burdens on business. The intended effect is that those businesses not already reducing the risks to their workers to a sufficiently low level take further, proportionate action to ensure that this is achieved.

What policy options have been considered? Please justify any preferred option.

Three policy options were considered: (1) do nothing - continue to rely on existing regulatory provisions; (2) rely on existing regulatory provisions where appropriate - introduce new regulatory provisions limited to new requirements set by the Directive where necessary and (3) introduce a full set of new regulatory provisions to reproduce the full requirements of the Directive disregarding existing regulatory provisions. Option 2 is preferred. Option 1 would not meet the Government's legal test for transposition; Option 3 would require unnecessary risk assessments to be undertaken by businesses which would not result in a reduced level of risk to workers and would not be in line with Better Regulation; Option 2 is considered the best fit to meet all 3 policy objectives.

When will the policy be reviewed to establish the actual costs and benefits and the achievement of the desired effects? 3 years from coming into force – April 2013

Ministerial Sign-off

For final proposals/implementation stage Impact Assessments:

I have read the Impact Assessment and I am satisfied that, given the available evidence, it represents a reasonable view of the likely costs, benefits and impact of the leading options.

Signed by the responsible Minister:

William D. McKenzie ................................................................. Date: 30 March 2010
**Summary: Analysis & Evidence**

**Policy Option: 1**

**Description:** Do nothing - continue to rely on existing regulatory provisions already in place on Great Britain

### Annual Costs

**One-off (Transition)**
- **Yrs:** N/a
- **£:** 0

**Average Annual Cost (excluding one-off)**
- **£:** 0
- **N/a**

**Total Cost (PV): £ 0**

**Other key non-monetised costs** by ‘main affected groups’

The fact that this option would not transpose the Directive would leave GB open to infraction proceedings from the European Commission. These are likely to require GB to develop new regulations – either Option 2 or 3 and may result in large financial penalties until delivered.

### Annual Benefits

**One-off**
- **Yrs:** N/a
- **£:** 0

**Average Annual Benefit (excluding one-off)**
- **£:** 0
- **N/a**

**Total Benefit (PV): £ 0**

**Other key non-monetised benefits** by ‘main affected groups’

N/a

### Key Assumptions/Sensitivities/Risks

N/a

### Price Base
- **Year:** N/a

### Time Period
- **Years:** 10

### Net Benefit Range (NPV)
- **£ 0**

### NET BENEFIT (NPV Best estimate)
- **£ 0**

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**What is the geographic coverage of the policy/option?**

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**On what date will the policy be implemented?**

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**Which organisation(s) will enforce the policy?**

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**What is the total annual cost of enforcement for these organisations?**

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<tbody>
<tr>
<td></td>
<td>£ N/a</td>
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**Does enforcement comply with Hampton principles?**

Yes

**Will implementation go beyond minimum EU requirements?**

No

**What is the value of the proposed offsetting measure per year?**

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<tr>
<td></td>
<td>£ N/a</td>
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**What is the value of changes in greenhouse gas emissions?**

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<tbody>
<tr>
<td></td>
<td>£ N/a</td>
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</table>

**Will the proposal have a significant impact on competition?**

No

**Annual cost (£-£) per organisation (excluding one-off)**

<table>
<thead>
<tr>
<th>Organisation Type</th>
<th>Micro</th>
<th>Small</th>
<th>Medium</th>
<th>Large</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N/a</td>
<td>N/a</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

**Are any of these organisations exempt?**

No

**Impact on Admin Burdens Baseline (2005 Prices)**

<table>
<thead>
<tr>
<th>Increase of</th>
<th>Decrease</th>
<th>Net Impact</th>
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<tbody>
<tr>
<td>£ 0</td>
<td>£ 0</td>
<td>£ 0</td>
</tr>
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</table>

**Key:** Annual costs and benefits: Constant
### Summary: Analysis & Evidence

| Policy Option: 2 | Description: Rely on existing regulatory provisions to introduce new provisions limited to specific relevant Directive where necessary |

### ANNUAL COSTS

<table>
<thead>
<tr>
<th>Description and scale of key monetised costs by ‘main affected groups’ (Some minor costs omitted)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Familiarisation: 1.1m to 1.7m, best = 1.4m</td>
</tr>
<tr>
<td>Worker information: 3m to 8.5m, best = 5.4m</td>
</tr>
<tr>
<td>Risk assessment: 1.6m to 4m, best = 2.7m</td>
</tr>
<tr>
<td>Reduce risk &amp; health surveillance: 1.6m to 4.7m, best = 2.9m</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Costs</th>
<th>Description and scale of key monetised costs by ‘main affected groups’</th>
</tr>
</thead>
<tbody>
<tr>
<td>One-off (Transition)</td>
<td>Yrs</td>
</tr>
<tr>
<td>Average Annual Cost (excluding one-off)</td>
<td>£ 1.1m (0.6 to 1.7m)</td>
</tr>
</tbody>
</table>

**Total Cost (PV)**: £ 12.6m (7.4 to 19.1m)

### ANNUAL BENEFITS

<table>
<thead>
<tr>
<th>Description and scale of key monetised benefits by ‘main affected groups’</th>
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<tbody>
<tr>
<td>Evidence suggests ill health due to AOR is low. As a result, very low levels of additional health benefits are expected. The maximum credible number of avoided cases was estimated to be 200 (top end of the range), with 2 taken as a minimum.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Benefits</th>
<th>Description and scale of key monetised benefits by ‘main affected groups’</th>
</tr>
</thead>
<tbody>
<tr>
<td>One-off</td>
<td>Yrs</td>
</tr>
<tr>
<td>Average Annual Benefit (excluding one-off)</td>
<td>£ 0m (0m to 0.1m)</td>
</tr>
</tbody>
</table>

**Total Benefit (PV)**: £ 0.3m (0m to 0.6m)

### Key Assumptions/Sensitivities/Risks

### Price Base

| Year 2009 | Time Period Years 10 | Net Benefit Range (NPV) | £ -19m to -6.8m | NET BENEFIT (NPV Best estimate) | £ -12.24 |

| What is the geographic coverage of the policy/option? | | |
|---|---|---|---|---|---|
| On what date will the policy be implemented? | | |
| Which organisation(s) will enforce the policy? | | |
| What is the total annual cost of enforcement for these organisations? | £ N/a | |
| Does enforcement comply with Hampton principles? | Yes | |
| Will implementation go beyond minimum EU requirements? | No | |
| What is the value of the proposed offsetting measure per year? | £ N/a | |
| What is the value of changes in greenhouse gas emissions? | £ N/a | |
| Will the proposal have a significant impact on competition? | No | |
| Annual cost (£-£) per organisation (excluding one-off) | Micro N/a | Small N/a | Medium N/a | Large N/a |
| Are any of these organisations exempt? | No | No | N/A | N/A |

**Impact on Admin Burdens Baseline (2005 Prices)** (Increase - Decrease)

| Increase of | £ 0 | Decrease | £ 0 | Net Impact | £ 0 |

**Kev:** Annual costs and (Net)
### Summary: Analysis & Evidence

| Policy Option: 3 | Description: Introduce a full set of new regulatory provisions to reproduce the full requirements of the Directive, disregarding

#### ANNUAL COSTS

<table>
<thead>
<tr>
<th>One-off (Transition)</th>
<th>Years</th>
<th>Description and scale of key monetised costs by ‘main affected groups’ (Some minor costs omitted)</th>
</tr>
</thead>
<tbody>
<tr>
<td>£5.2m (2.9 to 6.2m)</td>
<td>1</td>
<td>Familiarisation: 1.1m to 1.7m, best = 1.4m</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Worker information: 3m to 8.5m, best = 5.4m</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Risk assessment: 4.1m to 10.3m, best = 7.7m</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Reduce risk &amp; health surveillance: 1.7m to 5m, best = 3.1m</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Average Annual Cost (excluding one-off)</th>
<th>Years</th>
<th>Description and scale of key monetised costs by ‘main affected groups’</th>
</tr>
</thead>
<tbody>
<tr>
<td>£1.5m (0.8 to 2.3m)</td>
<td>10</td>
<td></td>
</tr>
</tbody>
</table>

| Total Cost (PV) | £17.9m (10.2 to 25.8m) |

#### ANNUAL BENEFITS

<table>
<thead>
<tr>
<th>One-off</th>
<th>Years</th>
<th>Description and scale of key monetised benefits by ‘main affected groups’</th>
</tr>
</thead>
<tbody>
<tr>
<td>£0</td>
<td>1</td>
<td>Benefits are expected to be the same as under Option 2 due to the low levels of baseline risk, and therefore opportunity to reduce risk through the additional risk assessment effort involved with Option 3.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Average Annual Benefit (excluding one-off)</th>
<th>Years</th>
<th>Description and scale of key monetised benefits by ‘main affected groups’</th>
</tr>
</thead>
<tbody>
<tr>
<td>£0m (0m to 0.1m)</td>
<td>10</td>
<td></td>
</tr>
</tbody>
</table>

| Total Benefit (PV) | £0.3m (0m to 0.6m) |

Other key non-monetised costs by ‘main affected groups’

#### Key Assumptions/Sensitivities/Risks

**Price Base**
- Year 2009

**Time Period**
- Years 10

<table>
<thead>
<tr>
<th>Net Benefit Range (NPV)</th>
<th>NET BENEFIT (NPV Best estimate)</th>
</tr>
</thead>
<tbody>
<tr>
<td>£-25.8m to -9.6m</td>
<td>£-17.6</td>
</tr>
</tbody>
</table>

What is the geographic coverage of the policy/option?

On what date will the policy be implemented?

Which organisation(s) will enforce the policy?

What is the total annual cost of enforcement for these organisations?

Does enforcement comply with Hampton principles?

Will implementation go beyond minimum EU requirements?

What is the value of the proposed offsetting measure per year?

What is the value of changes in greenhouse gas emissions?

Will the proposal have a significant impact on competition?

Annual cost (£-£) per organisation (excluding one-off)

<table>
<thead>
<tr>
<th>Micro N/a</th>
<th>Small N/a</th>
<th>Medium N/a</th>
<th>Large N/a</th>
</tr>
</thead>
</table>

Are any of these organisations exempt?

No

Impact on Admin Burdens Baseline (2005 Prices)

<table>
<thead>
<tr>
<th>Increase of</th>
<th>Decrease</th>
<th>Net Impact</th>
<th>Key:</th>
</tr>
</thead>
<tbody>
<tr>
<td>£0</td>
<td>£0</td>
<td>£0</td>
<td>Annual costs and benefits: Constant (Net Present)</td>
</tr>
</tbody>
</table>
Control of Artificial Optical Radiation at Work Regulations 2010

Aim of the proposal

2. To ensure that all workers in Great Britain are protected from hazardous sources of artificial optical radiation in the workplace and benefit from the requirements of the Physical Agents (Artificial Optical Radiation) Directive – ‘the Directive’.

Background

3. Optical radiation is another term for light; artificial sources of light (artificial optical radiation - AOR) in the workplace can generate visible, ultraviolet, infrared and laser radiation.

4. AOR is present in virtually all workplaces and the vast majority of sources pose no health and safety problems. However a minority of sources can produce sufficiently high levels of radiation to damage the eyes and/or skin of workers if they are not managed properly. For example ultraviolet radiation generated in welding can cause inflammation of the cornea (the condition ‘arc eye’) and laser radiation generated in a number of industrial and research processes can permanently damage eyes and skin.

5. These hazards are, in general, already well understood and well managed in Great Britain; inspectors do not come across many instances of workers at risk and there are very few cases of ill health or injury arising from known exposure to AOR reported.

6. Nevertheless, AOR hazards were considered sufficiently serious at a European level for the European Commission to propose a Directive to specify common control measures that need to be in place in workplaces across European Member States and for arrangements to be made to enforce these controls.

7. The Directive was adopted (‘approved’) in 2006¹ and must be transposed and implemented (its requirements brought into law) throughout the UK by 27 April 2010 to ensure a harmonised control regime across European Member States.

Reason for Government action

8. For the purposes of implementing this Directive, Great Britain, Northern Ireland and Gibraltar collectively make up the United Kingdom. The Health and Safety Executive (HSE) takes the lead for Government for ensuring the Directive’s requirements come into force in GB. This will be the focus of this impact assessment.

9. In considering the best method to achieve implementation, the policy objectives were to ensure that the eyes and skin of workers are protected from hazardous AOR sources in the workplace and that the Directive is implemented in a proportionate way which achieves the aims of the Directive while also taking into account existing controls and minimising unnecessary burdens on business.

10. Three different options were considered to meet these objectives:

- **Option 1**: Do nothing – continue to rely on existing regulatory provisions already in place in Great Britain
- **Option 2**: Rely on existing regulatory provisions where appropriate and introduce new regulatory provisions limited to new, specific requirements set by the Directive where necessary
- **Option 3**: Introduce a full set of new regulatory provisions to reproduce the full requirements of the Directive disregarding existing regulatory provisions.

The pros and cons of these different options are developed below.

11. **Option 1** proposes no change from the current situation. Evidence from a number of official data sources\(^2\) as well as inspector feedback indicates that the incidence of injury and ill health associated with AOR in the workplace is very rare. In the last 15 years there are estimated to have been fewer than 10 injuries that required workers to take more than 3 days off work reported to HSE and there have been no cases of work-related cataracts or neoplasia (new or abnormal tissue growth) attributed to AOR reported by general practitioners or occupational physicians. There have been 19 actual (30 estimated cases) of work related neoplasia reported by consultant dermatologists which were attributed by them to exposure to AOR through the Occupational Skin Surveillance Scheme (EPIDERM). In 90% of these cases, the workers were involved in welding which generates high levels of ultraviolet light. However, it was not clear how many of these workers also worked outside or spent their leisure time outside – which would also increase their exposure to natural optical radiation (sunlight). There have also been 65 cases of heat cataracts (a prescribed industrial disease) compensated under the Industrial Injuries scheme between 1992 and 2008. None of these occurred after 2002.

12. This indicates that, in general, AOR hazards are well understood and well managed. Welding and hot (eg foundry) work are traditional activities that have taken place in British workplaces for a great many years. This has allowed an awareness and appreciation of the risks to build up amongst workers, along with knowledge and adoption of sensible measures to manage the risks. It also reflects the valuable inputs from safety professionals and AOR specialists who have developed proportionate control measures for emerging AOR hazards, such as lasers.

13. In terms of the existing regulatory framework, there are no specific regulations for hazardous sources of AOR in Great Britain. However the Management of Health and Safety at Work Regulations 1999 (MHSWR) address the general principles of how hazards in the workplace need to be managed, through risk assessment and adoption of proportionate control measures to ensure that risks are reduced to as low a level as is reasonably practicable. The MHSWR are regularly used by businesses working with hazardous sources of AOR as a framework on which to develop a proportionate risk management system, as evidenced by the few reports of harm.

14. These risk management systems involve using control measures proportionate to the activity being undertaken. The Directive effectively codifies these thereby ensuring that they are in place across all Member States. However, because these measures are now listed in a Directive, it means that they must also be covered in national law. Existing health and safety legislation does not address these specific measures and as such cannot be relied on to transpose the Directive. As such Option 1 would not transpose the Directive in an appropriate way and is not considered further.

\(^2\) Reporting of Injuries, Diseases and Dangerous Occurrences Regulations 1995 (RIDDOR) and The Health and Occupation Reporting Network (THOR) which includes the discrete Occupational Skin Surveillance Scheme (EPIDERM), Occupational Physicians Reporting Activity (OPRA) and THOR-GP which covers General Practitioners)
15. **Option 2** represents a proportionate approach to achieve the aims of the Directive to protect the eyes and skin of workers. It builds on obligations already in place under existing regulations (eg MHSWR, Health and Safety (Consultation with Employees) Regulations) but includes new AOR-specific requirements where appropriate in order to ensure that businesses that need to take additional measures to reduce the level of risk associated with hazardous sources of AOR in their workplace do so.

16. In considering Option 2, HSE acknowledges that the vast majority of businesses do not need to do anything more to reduce the risks associated with AOR – either because they only have safe sources or already manage the risks associated with hazardous sources properly. Option 2 offers the opportunity to minimise unnecessary additional burdens by effectively removing these businesses from further obligations.

17. **Option 3** represents the more traditional approach to transpose European Health and Safety Directives, effectively reproducing all aspects of the Directive, even those that are already covered in existing health and safety law. One of the implications of this option is that all businesses would be required to undertake a potentially detailed risk assessment to determine whether the AOR sources they use are hazardous. Reproducing this exact requirement would make it more difficult to minimise unnecessary additional burdens, potentially placing responsibilities on dutyholders which we suspect will have little impact on the risks.

**Data sources and general assumptions**

18. In order to estimate the number of businesses potentially affected, HSE has used information from several sources:

   Work commissioned by HSE:
   - Review of occupational exposure to optical radiation and electric and magnetic fields with regard to the proposed CEC Physical Agents Directive, NRPB R265, 1994;

   European Commission information:

   UK information
   - Data supplied by EEF – The manufacturers’ organisation - on likely number of businesses in the manufacturing sector using hazardous sources of AOR and reasonable specialist health and safety consultant fees
   - Data obtained from the Office of National Statistics on the Annual Business Inquiry – workplace analysis – for the numbers of businesses likely to be undertaking the activities involving hazardous sources of AOR listed below.

19. Unless otherwise stated, all other assumptions are based on judgements applied by HSE’s technical specialists and feedback from stakeholders to the public consultation, which ran from November 2009 to February 2010. Costs have been discounted at an annual rate of 3.5% (in line with Treasury guidance). Prices are expressed in 2009 values.
Work activities likely to be affected by the Regulations

20. Activities involving hazardous sources of AOR that could pose a risk of harming the eyes and skin of workers include:

- Metal working – welding (both arc and oxy-fuel) and plasma cutting
- Pharmaceutical and research - UV fluorescence and sterilisation systems
- Hot industries – proximity to furnaces and molten products
- Printing – UV curing of inks
- Motor vehicle repairs – UV curing of paints (plus welding/cutting)
- Medical and cosmetic treatments – laser surgery, blue light and UV therapies, intense pulsed lights (IPLs)
- Research and education - all use of Class 3B and Class 4 lasers

Number of businesses likely to be affected by the Regulations

21. HSE interrogated the data sources listed above to estimate the numbers of businesses likely to be undertaking activities involving hazardous sources of AOR (for example we assumed that 100% of motor vehicle repair businesses will undertake welding and added this to 100% of businesses known to have furnaces).

22. HSE estimate the number of businesses using hazardous sources of AOR across all sectors to be 80,000.

Benefits

Health and safety benefits

23. Hazardous sources of AOR in the workplace can cause harm and need to be managed. However evidence collected (from RIDDOR, THOR and inspectors – see above) indicates that in Great Britain this hazard is already well understood where it occurs and well managed by individual businesses, industry groups and safety professionals and radiation specialists that deal directly with it, resulting in very few reported cases of ill health or injury.

24. This gives a very low baseline level of harm associated with AOR and as such we expect to see very limited direct health and safety benefits with either Option 2 or 3. Nevertheless, HSE is aware of anecdotal evidence of exposures resulting in short term, acute eye and/or skin conditions. For example, ’arc eye’ or ‘welders flash’ is a painful inflammation of the cornea, the clear tissue which covers the front of the eye, which can happen if the eye is exposed to an intense source of ultraviolet light, such as in types of welding. Although painful, the condition frequently clears after 1 or 2 days and rarely (if secondary infection occurs) results in long-term damage: as such, it is not reportable to HSE and as such may not appear in the statistics. In addition UV trans-illuminators used in molecular biology research have been known to cause short-term erythema (equivalent to sunburn) symptoms in researchers who have not taken appropriate precautions to protect their skin.

25. The fact that harm does still occur indicates that the risks could be managed more effectively in what we estimate to be a very small number of businesses. This clearly could result in realisation of some benefits for their workers.

26. The option chosen is likely therefore, to result in an increase in the level of protection offered to workers in the small proportion of businesses where risks are currently not being effectively managed, and where the employers respond appropriately to the
legislation. This will result in a reduction in the number of minor injuries attributable to AOR.

27. However, we are unable to estimate the numbers involved. Accepting that the cost to society of a minor injury is estimated at £350 we recognise that the economic value of health and safety benefits are likely to be outweighed by the costs in economic terms. This can be demonstrated by proposing a test of benefits scenarios in which the benefits of a maximum credible number of annual cases is calculated. Doing this shows that even with 200 avoided cases annually (this is anticipated to be pushing beyond what is credible given the baseline evidence) benefits to society would remain well below costs (£600k over a 10-year period). Alternative minimum and best estimate benefits estimates are made, but clearly do not change the overall message that monetised benefits are expected to be significantly lower than costs, under any credible cost scenario (minimum / best estimate / maximum cost).

Other benefits

28. There will also be unquantifiable benefits due to the harmonisation of control regimes across Member States with the Directive ensuring equity of worker protection. This may encourage freedom of movement of British workers, allowing them to work in other Member States under the same level of protection as in Great Britain.

29. There will also be minor benefits in competition terms to UK businesses with hazardous sources of AOR, as the Directive will provide greater harmonisation and more consistent control regimes in place in all businesses across Member States with hazardous sources of AOR.

Costs

Costs to Business

Option 1

30. There are no direct costs from this option. However, this will not transpose the Directive properly and is not a viable option for further scrutiny. Any ongoing costs and benefits of complying with the MHSWR will continue to be borne by businesses working with hazardous sources of AOR and using the MHSWR as a framework on which to develop a proportionate risk management system. There is no evaluative information on the ongoing costs of complying with the MHSWR for the relevant activities.

Option 2 & 3

31. These will be assessed together under common headings.

Cost of Screening

32. Businesses with employees that use hazardous sources of AOR will be in scope of new regulations developed under either option 2 or 3. In order to determine whether they are in scope of the new AOR Regulations we assume that a proportion of businesses (including those not intended to be in scope) will undertake a basic screen which we assume will involve a process similar to the flow charts at Annexes 1 and 2 and by looking at the list of safe and unsafe sources in the HSE guidance.

33. HSE will not be undertaking proactive communications around these new regulations. Furthermore, it is not immediately obvious to many employers what artificial optical radiation is and there is a low baseline of risk associated with this hazard. As a result of these facts, we assume that, for both options, only 3% (143,000) of total.

businesses in the UK will undertake screening and that they will spend 10 minutes looking at the guidance and flow chart at an estimated labour cost of £29.25 per hour (production manager - or equivalent). Many of these will be those who have hazardous sources of AOR, and are therefore more likely to recognise the term ‘artificial optical radiation’. We have a high level of confidence that a very low proportion of total UK businesses will screen. Increasing the 3% assumption to the boundary of what is considered likely, i.e. 5% does not have a significant impact on the total costs.

34. We therefore estimate the one-off costs of screening, for Option 2 and 3 will be around **£0.7 million** (best estimate with a range of £0.63m-0.77m).

### Cost of familiarisation and resolving uncertainty

35. Because of this screening, we assume that there will be a proportion of businesses that familiarise themselves to determine whether they need to do more. However, we assume that reading the HSE guidance will be sufficient for those businesses with obviously safe sources to identify that they do not need to take further action. Because there has already been considerable awareness raised amongst particular sectors about the forthcoming AOR Regulations, including through HSE’s public consultation, we assume that the proportion of the estimated 80,000 businesses with hazardous sources of AOR that will (a) be aware of them and (b) will need to undertake significant familiarisation with the regulations to investigate whether they need to do any form of further action will range from 20-40% (with a best estimate of 30%).

36. These businesses will range from small/medium sized enterprises through to large research and manufacturing enterprises. Within these businesses, there may only be a single activity involving hazardous AOR which is relatively straightforward in nature (e.g. welding) or there may be a number of different AOR activities of varying complexities (e.g. research involving lasers).

37. We assume that all the businesses will familiarise themselves and resolve uncertainty by studying the HSE guidance. They will identify that they use hazardous sources and are in scope. This will require varying levels of familiarisation, but we assume this will be the same under both option 2 and 3. Taking an average across all businesses and all activities, we assume that this will, on average, require 1 hour at an estimated labour cost of £29.25 per hour (production manager - or equivalent).

38. We therefore assume the first year, one-off costs of familiarisation by those businesses with hazardous sources for Option 2 and 3 will be around **£0.7 million** (best estimate with a range of £0.5m-0.9 m).

### Cost of refreshing existing risk assessments and addressing uncertainty

39. It is from risk assessment onwards that the costs associated with Options 2 and 3 start to diverge.

#### Option 2

40. Option 2 would facilitate greater flexibility, requiring additional risk assessment activity to be undertaken only when (a) work involves hazardous sources of AOR that could harm the eyes and/or skin **AND** (b) measures have not already been implemented which reduce the risk to as low a level as is reasonably practicable.

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4 http://stats.berr.gov.uk/ed/sme/smestats2008.xls. 4.8 million companies in UK

5 Source of information: Annual Survey of Hours and Earnings, earnings for production manager have been used and adjusted by a factor of 1.3 to convert earnings into total costs (to include overheads etc). http://www.statistics.gov.uk/statBase/product.asp?vlnk=13101
41. We already have evidence of a very low level of employee exposure to risk within the baseline for this impact assessment, indicating that the majority of existing risk assessments are suitable and sufficient.

42. Nevertheless, we recognise that a proportion of businesses that already expose their employees to an appropriately low level of risk will take the opportunity to refresh their risk assessments. Whilst this will not give any benefits in terms of further reductions in risk, it will give unquantifiable reassurance that they do not need to take any further action. The costs associated with refreshing risk assessments are estimated below.

43. We have already assumed that 20-40% of the 80,000 businesses with hazardous sources of AOR will familiarise themselves with the AOR Regulations.

44. We now assume that 40% of these will do no more and that 30% of these will refresh their risk assessments even though their staff are at an acceptably low risk.

45. We assume that this will be achieved by cross-referencing their existing arrangements with the control measures outlined in the HSE guidance, and updating where appropriate.

46. In total, this may result in more than one risk assessment being refreshed, but we assume that each affected business will, on average, spend 2 hours refreshing their risk assessments at an estimated labour cost of £29.25 per hour.

47. We therefore estimate the one-off costs for refreshing risk assessments and resolving uncertainty for Option 2 will be £0.4 million (best estimate with a range of £0.3m-0.6 m).

Option 2

48. Option 2 would require all businesses with hazardous sources to develop a new risk assessment; as such a simple refreshing of existing risk assessments would not be appropriate.

49. We therefore estimate there will be no one-off costs for refreshing risk assessments and resolving uncertainty for Option 3.

Cost of developing new risk assessments to reduce risks

50. We have already assumed that of the 20-40% of businesses with hazardous sources of AOR who have familiarised themselves with the new regulations, 40% will do no further work on their risk assessments and 30% will refresh them.

51. We now assume that the remaining 30% will develop a new risk, AOR-specific risk assessment. As this risk assessment is new, it involves a one off effort up front but also recurring activity to maintain risk assessments. Businesses are assumed to update their risk assessments approximately every 3 years and mirror the original effort, and are in line with costs of refreshing risk assessments outlined in the previous section. On that basis, we assume both one off first year, and recurring costs over the appraisal period (10 years) in order to calculate a total present value cost.

52. In calculating the one off costs for option 2, we assume that 70% of these will follow the HSE guidance, sector specific guidance and use information provided by manufacturers and revise the risk assessments themselves. This may result in more than one risk assessment being revised but we assume that each affected business will, on average, spend 3 hours reading the HSE guidance and revising their risk assessments at an estimated labour cost of £29.25 per hour.
53. We recognise that the specific risk assessment requirements are new, prescriptive and potentially complex and that there is likely to be a lack of in-house expertise or competence, particularly where measurements are being considered. We therefore assume that 30% of businesses developing new risk assessments will use a specialist or consultant. We assume that these will cost on average £750 per day and will on average spend 0.5 days revising the risk assessment.

54. We therefore estimate the total one off costs for developing new risk assessments under Option 2 will be £1.25 million (best estimate with a range of £0.7-2.0 million).

55. Recurring costs are calculated on the basis that all businesses that undertook a new risk assessment would update typically every three years, involving around 2 hours of a manager’s time at the same labour cost. This results in a cost, recurring every 3 years, with a present value over ten years of around £1.03 million (best estimate with a range of £0.6m-1.5 m).

56. Total one off and recurring costs associated with new risk assessment under Option 2 results in a best estimate present value of £2.3 million (with a range of £1.3m-3.4m).

Option 3

57. We have already assumed that of the 20-40% of businesses with hazardous sources of AOR will familiarise themselves with the new regulations. For option 2 we have assumed 40% of these will do no further work, 30% will refresh their risk assessment and 30% will develop new risk assessments.

58. Option 3 requires more detailed revisions of risks assessment given that refreshing risk assessments would not be appropriate. We therefore assume for that, of those businesses that familiarise themselves with the new regulations, 40% will do no further work and 60% will develop new risk assessments. In addition, the simple guidance HSE would develop for Option 2 is unlikely to be sufficiently detailed to enable dutyholders to comply with their obligations and further reading or input from specialists will be required. As such, some of the assumptions will differ between Options 2 and 3.

59. In calculating the one off costs for option 3, we assume that 50% of these will follow the HSE guidance, European guidance, sector specific guidance and use information provided by manufacturers and revise the risk assessments themselves. This may result in more than one risk assessment being revised but we assume that each affected business will, on average, spend 4 hours revising their risk assessments at an estimated labour cost of £29.25 per hour.

60. We also assume that 50% of businesses developing new risk assessments under Option 3 will use a specialist or consultant. We assume that these will cost on average £750 per day and will on average spend 0.5 days revising the risk assessment.

61. We therefore estimate the total one off costs for developing new risk assessments under Option 3 will be £3.5 million (best estimate with a range of £1.6m to 4.3m).

62. Recurring costs are calculated on the basis that all businesses that undertook a new risk assessment would update typically every three years, involving around 4 hours of a manager’s time at the same labour cost. This results in a cost of around £1.7 million (best estimate with a range of £1 to 2.5 million), recurring every 3 years, with a present value over ten years of around £4.1 million (best estimate with a range of £2.5 to 6 million).

63. Total one off and recurring costs associated with new risk assessment under Option 3 results in a best estimate present value of £7.7 million (with a range of £4m to 10 m).
Cost of reducing risks

64. We have already assumed a low baseline level of risk associated with this hazard. Nevertheless, whichever option is selected would need to ensure that the minority of businesses where the risks are not already reduced to a sufficiently low level adopt control measures to achieve this.

Option 2

65. Because of the low baseline level of risk, we are confident that only a small proportion of those businesses that revised their risk assessment will identify the need to take further measures to reduce the AOR risks to their employees. We therefore assume this number to be 10% (720).

66. To identify the controls needed, they will either use available guidance or will employ a specialist or consultant. The changes we expect to see made will involve a combination of organisational factors (such as demarcating areas where hazardous sources of AOR are used) and hardware issues (such as buying new PPE which is better suited to the activity and/or replacing old pieces of equipment for ones that better protect against AOR at source).

67. We have divided the impact costs into two components – the time taken to implement the changes and the cost of any equipment. In calculating the time, we assume that 70% of businesses will make the changes themselves and 30% will use consultants. Those undertaking themselves will follow the HSE guidance, sector specific guidance and use information provided by manufacturers and each affected business will, on average, spend 2 hours making the changes at an estimated labour cost of £29.25 per hour. Those employing consultants, we assume will cost on average £750 per day and will on average spend 0.5 days.

68. Recurring costs associated with time are calculated on the basis that all affected businesses will review their control measures every three years with the time and labour costs being the same as above. This results in a cost, recurring every 3 years, with a present value over ten years of around £0.38 million (best estimate with a range of £0.2-0.64 million).

69. Not all businesses will require new equipment. Averaging out the cost of new equipment over the 720 businesses affected, we assume will cost, on average, £1000 per business. This is assumed to be a recurring cost, which recurs typically every 3 years. This produces a best estimate present value over ten years of £2.48 million (with a range of £1.34m-4.01 million).

70. Total one off and recurring costs associated with implementing new control measures to reduce AOR risks under Option 2 results in a best estimate present value of £2.86 million (with a range of £1.54m-4.65 million).

Option 3

71. We assume that the same amount of businesses will identify that they need to take further measures whether they undertook risk assessments under Option 2 or Option 3. We therefore assume this number to be 720.

72. Businesses adopting control measures under Option 3 must develop an action plan as specified in the Directive. This will include a prescriptive list of measures required to reduce worker exposure below the exposure limit values.

73. Because the assessment of exposure limit values is technically complex, we assume that a larger proportion of businesses will use consultants, and the task itself is more time intensive than under Option 2. Specifically, 50% of businesses affected will develop an action plan themselves, following European guidance, sector specific guidance and undertaking measurements. We assume that each affected business...
will, on average, spend 4 hours developing and implementing an action plan at an estimated labour cost of £29.25 per hour.

74. The remaining 50% will employ consultants at a cost on average of £750 per day and who will on average spend 0.5 days on the action plans.

75. Recurring costs associated with time are calculated on the basis that all affected businesses will review their control measures every three years with the time and labour costs being the same as above. This results in a cost, recurring every 3 years, with a present value over ten years of around £0.6 million (best estimate with a range of £0.4-0.9 million).

76. We assume that the action plans developed under option 3 will not result in any changes to the typical equipment costs assessed under option 2. Equipment costs for option 3 are therefore assumed the same as for Option 2.

77. Total one off and recurring costs associated with implementing new control measures to reduce AOR risks under Option 3 results in a best estimate present value of £3 million (with a range of £1.7-5.0 million).

Cost of providing information and training

Option 2 & 3

78. We assume that 100% (7200) of those businesses that develop new risk assessments will deliver additional training to their staff to ensure they understand what control measures and working practices they need to adopt to reduce the AOR risks. This training is likely to have a big impact on risk reduction by increasing the likelihood of staff adhering to local rules. Given that this number is the same for Option 2 or 3, we will assume the information and training costs will be the same for each option.

79. We assume that 20 – 30% of staff in each affected business will require additional training of 30 minutes at an average labour cost of £18 per hour. We assume that this training will be delivered by production managers or safety professionals (or equivalent) time at an average labour cost of £29.25 per hour. We assume that, in order to develop and deliver this training, the trainers themselves will need to be trained so that they can continue to provide advice on AOR hazards. Taking an average across all businesses affected, we assume, on average, trainers will take 90 minutes to familiarise themselves with the topic and to develop the training and will then take 30 minutes to deliver.

80. We assume that this training will recur on an annual basis requiring the same preparation and delivery times, with an annual cost for both trainers and trainees of around £0.6 million (best estimate with a range of £0.4m-0.10 million). Total annual costs associated with providing information and training under either Option 2 or 3 results in a best estimate 10 year present value of £5.45 million (with a range of £3.1m-8.5m).

Cost of providing medical examinations and appropriate health surveillance

Option 2 & 3

81. The requirement to provide medical examination and support in the event of an accidental overexposure to AOR, along with appropriate health surveillance is already enshrined in the Management of Health and Safety at Work Regulations 1999.

82. There will be no requirement for routine eye examinations as part of the final regulations developed under either Option 2 or 3. This is on the basis that the guidance published by the European Commission to accompany the Directive states

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6 The UK average hourly wage, from the Annual Survey of Hours and Earnings (ASHE) 2009, plus an extra 30% for non-wage labour costs.
that ‘50 years of experience has shown that such examinations have no value as part of a health surveillance programme and possibly introduce an additional risk to the worker…. A worker exposed to artificial optical radiation at work should not receive pre-employment, routine and post-employment eye examinations, just because they carry out such work’.

83. However the final regulations will require a worker whose eyes have been accidentally over-exposed to be offered a medical examination. Given that this mirrors current MHSWR requirements, this will place no additional duties on employers and therefore no additional cost.

84. Regulations developed under either Option 2 or 3 will place a specific requirement on employers to consider including skin surveillance in the event of a known over exposure. The guidance produced by the Commission states that "skin examinations are not usually justified purely on the basis of routine exposure to artificial optical radiation." We therefore interpret this to mean that examinations will only be required where the skin has been exposed to high levels of AOR, for example as the result of an accident, and because there are skin tests that occupational health providers could undertake, we will include this in the regulations.

85. As for accidental over-exposure of the eyes, workers whose skin has been over-exposed must be offered a medical examination, but given that this mirrors current MHSWR requirements, this will place no additional duties and therefore no cost.

86. Any additional skin surveillance will be undertaken at the direction of an occupational health provider but could involve examinations of the area of the skin known to have been subject to over-exposure for any changes which could be linked to exposure to high levels of AOR coupled with, for example, a self-reporting system on what changes in the skin to look for.

87. Whilst these will be additional requirements, because of the low baseline level of risk associated with this hazard, accidental over exposures are already considered to be very rare and will be even more so as a result of the new regulations. Assuming that 5 overexposures will still occur each year, with an annual cost for health surveillance of £1000 per business affected, this will result in a present value of around £43,000 over 10 years (best estimate with range of £35k-52k).

Additional costs not already covered.

88. We assume that there will be no additional costs associated with insurance premiums for either Option 2 or 3.

Costs to HSE

89. HSE envisages no change to its enforcement strategy when the AOR regulations come into force. The main costs to fall on HSE will be in the development and maintenance of guidance. This will be easier to do for Option 2 and will be met through HSE’s internal resources. Assuming the completion of the guidance requires the equivalent of three experts working full time on the guidance over a six month period, at an average gross wage rate of £40k7 this would amount to around £98,000.

90. Guidance to support Option 3 is less likely to be delivered within HSE given its technical nature and this may need to be developed via an external contract. Assuming this requires around 300 days of consultants' time at a day rate of around £750, this would cost HSE around £225,000.

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7 Adjusted upwards to account for overheads by a factor of 1.3.
Total costs to society

91. The total cost to society will consist of two main components: the cost to employers of complying with the new requirements, and the cost to the HSE of implementing and enforcing them. The costs associated with the Option 2 and Option 3 are summarised in the tables below.

92. Total one off and recurring costs associated with implementing Option 2 results in a best estimate present value of £12.51 million (with a range of £7.35m - £19.01m).

93. Total one off and recurring costs associated with implementing Option 3 results in a best estimate present value of £17.83 million (with a range of £10.12m - £25.75m).

94. Taking into account the costs, the better regulation agenda and feedback from stakeholders, Option 2 was considered to be the best fit for the policy objectives. A summary of the Option 2 compliance assumptions and headline costs is provided in Appendix 3.

95. A statement by HSE’s chief economist is provided at Appendix 4.
### Table 1: Summary of Costs associated with Option 2

<table>
<thead>
<tr>
<th></th>
<th>First year costs</th>
<th>Ten year present value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>£ million</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>Best estimate</td>
</tr>
<tr>
<td><strong>Implementation costs</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Self screening</td>
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<td>0.70</td>
</tr>
<tr>
<td>Familiarisation</td>
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<tr>
<td>Worker information</td>
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<td>0.63</td>
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<tr>
<td><strong>Policy Costs</strong></td>
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<td>Refresh risk assessment</td>
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<td>New risk assessment (one off)</td>
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<tr>
<td>New risk assessment (recurring)</td>
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<td>0.00</td>
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<tr>
<td>Action to reduce exposure (time)</td>
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<tr>
<td>Action to reduce exposure (equipment)</td>
<td>0.39</td>
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<tr>
<td>Health surveillance</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td><strong>HSE costs</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Produce guidance</td>
<td>0.09</td>
<td>0.10</td>
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<tr>
<td><strong>TOTAL</strong></td>
<td>2.96</td>
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Table 2: Summary of Costs associated with Option 3

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<th>Cost Category</th>
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<th>Ten year present value</th>
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<td>Min</td>
<td>Best estimate</td>
</tr>
<tr>
<td>Implementation costs</td>
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<td></td>
</tr>
<tr>
<td>Self screening</td>
<td>0.63</td>
<td>0.70</td>
</tr>
<tr>
<td>Familiarisation</td>
<td>0.47</td>
<td>0.70</td>
</tr>
<tr>
<td>Worker information</td>
<td>0.35</td>
<td>0.63</td>
</tr>
<tr>
<td>Policy Costs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Refresh risk assessment</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>New risk assessment (one off)</td>
<td>1.61</td>
<td>3.54</td>
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<tr>
<td>New risk assessment (recurring)</td>
<td>0.00</td>
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<td>Action to reduce exposure (time)</td>
<td>0.10</td>
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<tr>
<td>Action to reduce exposure (equipment)</td>
<td>0.39</td>
<td>0.72</td>
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<tr>
<td>Health surveillance</td>
<td>0.00</td>
<td>0.01</td>
</tr>
<tr>
<td>HSE costs</td>
<td></td>
<td></td>
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<tr>
<td>Produce guidance</td>
<td>0.20</td>
<td>0.23</td>
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<tr>
<td>TOTAL</td>
<td>3.75</td>
<td>6.70</td>
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</table>
Specific Impact Tests

Statutory Equality duties

96. No negative impact on equality on any of the groups addressed are expected. Hazardous sources of AOR are used in a variety of work activities undertaken by all groups covered by equality aspects and for which the control measures prescribed should give a comparable level of protection. However we will monitor relevant measurable outcomes to determine whether any group is detrimentally affected.

Economic Impacts: Competition

97. The Directive is being implemented across European Member States. As such the AOR regulations will reduce the potential for competitive advantage from lower costs/standards associated with control of AOR hazards. This will contribute towards a more level playing field in the EU and therefore is, in general terms, likely to have a positive impact on competition for UK businesses, which already have high level of protection.

98. The Directive represents, for the majority of UK employers, an additional cost versus very low marginal benefits, due to a high level of protection already in place. This represents some additional costs for employers, but it is not expected to have significant impacts on the number or range of suppliers, or significantly limit their ability to compete.

Economic Impacts: Small Firms

99. The majority of the 80,000 businesses using hazardous sources of AOR will be SMEs, in particular those undertaking welding. As such it is likely that SMEs will be impacted on more than other business types as a result of these regulations.
Use the table below to demonstrate how broadly you have considered the potential impacts of your policy options.

Ensure that the results of any tests that impact on the cost-benefit analysis are contained within the main evidence base; other results may be annexed.

<table>
<thead>
<tr>
<th>Type of testing undertaken</th>
<th>Results in Evidence Base?</th>
<th>Results annexed?</th>
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<tbody>
<tr>
<td>Competition Assessment</td>
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<td>No</td>
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<tr>
<td>Small Firms Impact Test</td>
<td>Yes</td>
<td>No</td>
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<tr>
<td>Legal Aid</td>
<td>Yes</td>
<td>No</td>
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<tr>
<td>Sustainable Development</td>
<td>Yes</td>
<td>No</td>
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<tr>
<td>Carbon Assessment</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Other Environment</td>
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<tr>
<td>Health Impact Assessment</td>
<td>Yes</td>
<td>No</td>
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<tr>
<td>Race Equality</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Disability Equality</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Gender Equality</td>
<td>Yes</td>
<td>No</td>
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<tr>
<td>Human Rights</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Rural Proofing</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>
Annex 1 - flow chart to be used by businesses who familiarise themselves with new AOR Regulations developed under Option 2

Start here: Main column

- **APPLICATION**
  - Are you an employer in Great Britain?
    - Yes
    - No
    - Unsure

- **SCOPE**
  - Do you use hazardous sources of Artificial Optical Radiation?
    - Yes
    - No
    - Unsure

- **RISK ASSESSMENT**
  - Are any risks to the eyes and skin already reduced to as low as reasonably practicable?
    - Yes
    - No
    - Unsure
  - A new risk assessment will be required

- **CONTROL RISKS**
  - Are any risks to eyes and skin reduced to as low as reasonably practicable
    - Yes
    - No

- **INFORMATION & TRAINING**
  - Develop and implement action plan to reduce risks
  - If new risk assessment indicates eyes and skin of workers are at risk, provide information and training on measures needed

- **MEDICAL EXAM & HEALTH SURVEILLANCE**
  - If accident/over-exposure, must ensure medical examination made available and appropriate health surveillance in place
    - Involve workforce
    - Involve occupational health provider as appropriate

- **COMPLIANT**
  - No Further Action Needed
Annex 2 - flow chart to be used by businesses who familiarise themselves with new AOR Regulations developed under Option 3

Start here: Main column

**APPLICATION**

Are you an employer in Great Britain?

**SCOPE**

Do you use hazardous sources of Artificial Optical Radiation?

- Yes
- Unsure
- No

A new risk assessment will be required

**RISK ASSESSMENT**

Develop and implement action plan to reduce risks

**CONTROL RISKS**

If new risk assessment indicates eyes and skin of workers are at risk, provide information and training on measures needed

**INFORMATION & TRAINING**

If accident/over-exposure, must ensure medical examination made available and appropriate health surveillance in place

**MEDICAL EXAM & HEALTH SURVEILLANCE**

Involve workforce

Involve occupational health provider as appropriate

No Further Action Needed

Look at HSE guidance or sector specific guidance

Use HSE & sector specific, EU guidance and manufacturers data. Involve workforce and competent adviser
### Annex 3: Summary of compliance assumptions associated with the costs of Option 2

<table>
<thead>
<tr>
<th>Activity</th>
<th>Businesses Affected</th>
<th>Time or Equipment costs</th>
<th>Best estimate of costs (10 year present value) £million</th>
</tr>
</thead>
<tbody>
<tr>
<td>Screening</td>
<td>3% of 1.2 million with employees 143,000</td>
<td>10 mins @ £29.25 per hour</td>
<td>0.7</td>
</tr>
<tr>
<td>Familiarisation</td>
<td>30% of 80,000 with hazardous sources 24,000</td>
<td>1 hour @ £29.25 per hour</td>
<td>0.7</td>
</tr>
<tr>
<td>Risk assessments – refresh</td>
<td>30% of 24,000 who familiarise 7,200</td>
<td>2 hours @ £29.25 per hour</td>
<td>0.4</td>
</tr>
<tr>
<td>Risk assessment – new</td>
<td>30% of 24,000 who familiarise 7,200</td>
<td>70% for 3 hours @ £29.25 per hour 30% use consultants for 0.5 day @ £750 per day 2.3</td>
<td></td>
</tr>
<tr>
<td>Control risks – time + equipment</td>
<td>10% of 7,200 who develop new RA 720</td>
<td>70% for 2 hours @ £29.25 per hour 30% use consultants for 0.5 day @ £750 per day 720 x £1000 for new equipment 2.86</td>
<td></td>
</tr>
<tr>
<td>Information and training</td>
<td>100% of 7,200 who develop new RA 7,200</td>
<td>7200 trainers for 1.5 hours @ £29.25 per hour 23,400 trainees for 0.5 hours @ £18 per hour 5.45</td>
<td></td>
</tr>
<tr>
<td>Medical Examination and Health surveillance</td>
<td>-</td>
<td>5 per year</td>
<td>£1000 per incident 0.04</td>
</tr>
<tr>
<td>HSE guidance</td>
<td>-</td>
<td>-</td>
<td>- 0.1</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
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<td><strong>12.55</strong></td>
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</table>
Annex 4: Statement by Chief Economist, Health and Safety Executive

As HSE Chief Economist I confirm that the attached Impact Assessment (IA), prepared by HSE Work Environment, Radiation & Gas Division in collaboration with the Economic Analysis Unit, makes appropriate use of evidence in analysing the costs and benefits of the alternative options.

The Control of Artificial Optical Radiation (AOR) at Work Regulations 2010 are designed to transpose into Great Britain law the requirements of the European Directive 'Physical Agents (Artificial Optical Radiation)', so ensuring that the risks to workers from AOR (e.g. ultraviolet radiation) are as low as reasonably practicable, while minimising the regulatory burden on business.

The IA considers three options: (1) relying on existing regulations (i.e. 'do nothing'); (2) relying on existing regulations but introducing new provisions aimed specifically at those businesses where risks associated with hazardous sources of AOR are not already adequately controlled; and (3) introducing a new full set of regulations to reproduce all aspects of the Directive, even those already covered in existing health and safety law.

The IA reviews the evidence from various statistical sources and concludes that the baseline level of injuries and ill health in Great Britain from exposure to AOR is very low. This has important implications for the assessment of both benefits and costs.

The health and safety benefits of introducing new regulations (Options 2 or 3) are expected to be limited: the prevention of a small number of minor injuries. Based on HSE estimates of the cost to society of such injuries, and of the maximum credible number that could be prevented, the total benefits discounted over a ten-year appraisal period are estimated to be less than £0.6 million. There would be some further unquantified benefits in terms of EU harmonisation and competition. The benefits are not expected to differ significantly between Options 2 and 3.

Some of the costs are also estimated to be the same for both Options 2 and 3: businesses' familiarisation with the new Regulations including 'screening' to see if they are affected (best estimate of costs over ten years = £1.4 million), and provision of information and training to workers (£5.4 million). The main difference between the options relates to the development of risk assessments, some of which will require the use of consultants: Option 2 would involve some businesses 'refreshing' their existing risk assessments (best estimate of ten-year costs = £0.4 million) and some developing new ones (£2.3 million), while Option 3 would involve no 'refreshing' but significantly more new assessments (£7.7 million). The costs of actions to reduce exposure, and of health surveillance, would be relatively low, because of the low level of baseline risk. These too would be a little higher under Option 3 (£3.1 million compared with £2.9 million under Option 2), as would the costs to HSE of producing guidance (£0.2 million compared with 0.1 million). The costs of Option 1 have not been quantified; however a significant non-monetised cost for this option would be the risk of being subject to infraction proceedings from
the European Commission, which would involve major financial penalties as well as legal costs and reputational damage.

The IA makes a number of assumptions, based on a mixture of statistical data and informed judgement, about the number of businesses affected and the proportions that will take certain actions. Where appropriate, minimum and maximum figures are given to indicate the range of uncertainty, and sensitivity analyses have been performed to check that varying the assumptions within credible limits would not alter the broad conclusions. Nonetheless there must remain considerable doubt as to the scale of the likely benefits and costs.

The IA’s preferred option is Option 2: this has significantly lower total estimated costs to society than Option 3 (by £5.3 million, or 30 per cent, over a ten-year appraisal period), without significantly lower expected benefits, and is preferable on grounds of proportionality and better regulation. Given that Option 1 has been ruled out because it would not transpose the Directive appropriately, I am satisfied that the evidence supports the choice of Option 2.

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Health & Safety Executive
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0151 951 4556
alan.spence@hse.gsi.gov.uk