



# **Cervical Cancer**

## **Measurability of Quality Performance Indicators Version 1.2**

**To be read in conjunction with:**

- **Cervical Cancer QPIs Final Publication v1.1**
- **Cervical QPI Dataset (Latest published version (v1))**

## Measurability of Quality Performance Indicators for Cervical Cancer

### Document control:

#### This Version

<b>Title</b>	Cervical Cancer Measurability of QPIs	<b>Version/Issue Number</b>	1.2
<b>Effective From</b>	October 2014	<b>Author</b>	Charlotte Anthony, ISD
<b>Document Type</b>	Guidance	<b>Document status</b>	Final – For Publication
<b>Document Purpose</b>	Final – For Publication		
<b>Summary of changes</b>			

#### Revision History

Version	Date	Status	Summary of Changes	QPI (s)
0.1	Feb 2014	Draft	First draft	All
0.2	Mar 2014	Draft	Changes following comments from KC	All
0.3	May 2014	Draft	Changes following public engagement	All
0.4	June 2014	Draft	Further changes following public engagement	All
0.5	Sept 2014	Draft	Changes following board review	All
1.1	July 2015	Final	Changes agreed at 9mth Review and back dated	2, 3, 6,7
1.2	April 2016	Final	Amendments outwith review	1

#### Updates from Previous Version

QPI	Summary of changes (excluding formatting changes) (April 2016)
1	Numerator changed from ([MRIDATE <> 10/10/1010] AND [MRIDATE < FIRSTTREATDATE]) to ([MRIDATE <> 10/10/1010] AND [MRIDATE ≤ FIRSTTREATDATE])

QPI	Summary of changes (excluding formatting changes) (July 2015)
1	Remove abdomen from description and numerator
2	Den – add OR 06
3	Num - [MDTDATE <> 10/10/1010] AND ([MDTDATE ≤ DEFTREATDATE] OR [DEFTREATDATE = 10/10/1010])
6	NR numerator - remove [RADIOTYPE = 99] OR
7	Num - add OR CHEMTYPE2 =05

## QPI 1 – Radiological Staging

<b>QPI Title:</b>	Patients with cervical cancer should have their stage of disease assessed by magnetic resonance imaging (MRI) prior to first treatment.
<b>Description:</b>	Proportion of patients with cervical cancer who have an MRI of the pelvis performed prior to first treatment.
Numerator	<p>Number of patients with cervical cancer having MRI of the pelvis carried out prior to first treatment.</p> <p>Date of MRI Scan Completed (Pre-treatment) not coded as not applicable and coded less than or equal to Date of First Treatment.</p> <p>([MRIDATE &lt;&gt; 10/10/1010] AND [MRIDATE ≤ FIRSTTREATDATE])</p>
Denominator	<p>All patients with cervical cancer. (Excluding patients with histopathological FIGO<sup>a</sup> stage IA1 disease, patients treated by LLETZ<sup>b</sup> only and patients unable to undergo MRI due to contraindications).</p> <p>Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; Final FIGO Stage {Cervical Cancer} not coded as IA1; and Surgery Performed not coded as LEEP/LLETZ (LEEP, LLETZ OR CONE); and MRI Scan {Pre-Treatment} not coded as Contraindication to MRI.</p> <p>([FIGO &lt;&gt; IA1] AND [SURG &lt;&gt;05 OR 06])) AND [MRI &lt;&gt; 98]</p>
Not recorded for numerator	<p>Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target</p> <p>[MRIDATE = 09/09/0909] OR [FIRSTTREATDATE = 09/09/0909]</p>
Not recorded for exclusion	<p>Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.</p> <p>[FIGO = 99] OR [SURG = 99] OR [MRI = 99]</p>
Not recorded for denominator	<p>Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard</p>

<sup>a</sup> FIGO – International Federation of Gynecology and Obstetrics

<sup>b</sup> Large Loop Excision of the Transformation Zone

## QPI 2 – Positron Emission Tomography/Computed Tomography (PET/CT)

<b>QPI Title:</b>	Patients with cervical cancer, for whom primary definitive surgery is not appropriate, should undergo positron emission tomography - computed tomography imaging (PET/CT).
<b>Description:</b>	Proportion of patients with cervical cancer, for whom primary definitive treatment is radical radiotherapy, who have PET/CT imaging.
Numerator	<p>Number of cervical cancer patients undergoing primary radical radiotherapy who have PET/CT imaging prior to starting treatment.</p> <p>Date of PET CT Scan {Cervical Cancer} (Pre-treatment) not coded as not applicable and coded as before Date Treatment Started (Radiotherapy)</p> <p>[PETSCAN &lt;&gt; 10/10/1010] <b>AND</b> [PETSCAN &lt; RSTARTDATE]</p>
Denominator	<p>All patients with cervical cancer undergoing primary radical radiotherapy. <i>(No exclusions)</i>.</p> <p>Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; Radiotherapy Course Type coded as Radical OR Chemoradiotherapy.</p> <p>([RADIOTYPE = 02 OR 06])</p>
Not recorded for numerator	<p>Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target</p> <p>[PETSCAN = 09/09/0909] <b>OR</b> [RSTARTDATE = 09/09/0909]</p>
Not recorded for exclusion	<p>Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.</p>
Not recorded for denominator	<p>Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard</p> <p>[RADIOTYPE = 99]</p>

### QPI 3 – Multidisciplinary Team Meeting (MDT)

<b>QPI Title:</b>	Patients with cervical cancer should be discussed by a multidisciplinary team (MDT) prior to definitive treatment.
<b>Description:</b>	Proportion of patients with cervical cancer who are discussed at a MDT meeting before definitive treatment.
Numerator	Number of patients with cervical cancer discussed at the MDT before definitive treatment.  Date discussed by Care Team (MDT) not coded as Not applicable and Date discussed by Care Team (MDT) before or equal to Date of Definitive Treatment {Cervical Cancer} OR Date of Definitive Treatment {Cervical Cancer} coded as not applicable [MDTDATE <> 10/10/1010] AND ([MDTDATE ≤ DEFTREATDATE] OR [DEFTREATDATE = 10/10/1010])
Denominator	All patients with cervical cancer. <i>(Excluding patients with histopathological FIGO<sup>c</sup> stage IA1 disease and patients treated by LLETZ<sup>d</sup> only).</i>  Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; and Final FIGO Stage {Cervical Cancer} not coded as IA1 and Surgery Performed not coded as LEEP/LLETZ(CONE) only.  ([FIGO <> IA1] AND [SURG <>05 OR 06])
Not recorded for numerator	Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target  [MDTDATE = 09/09/0909] OR [DEFTREATDATE = 09/09/0909]
Not recorded for exclusion	Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.  [FIGO = 99] OR [SURG = 99]
Not recorded for denominator	Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard

<sup>c</sup> FIGO – International Federation of Gynecology and Obstetrics

<sup>d</sup> Large Loop Excision of the Transformation Zone

## QPI 4 – Radical Hysterectomy

<b>QPI Title:</b>	Patients with stage IB1 cervical cancer should undergo radical hysterectomy.
<b>Description:</b>	Proportion of patients with stage IB1 cervical cancer (as defined by radiology and/or histopathology) who undergo radical hysterectomy.
Numerator	Number of patients with FIGO stage IB1 cervical cancer who undergo radical hysterectomy.  Surgery Performed coded as Radical Hysterectomy.  [SURG = 02]
Denominator	All patients with FIGO stage IB1 cervical cancer. <i>(Excluding patients who decline surgery, patients who undergo fertility conserving surgery and patients having neo adjuvant chemotherapy, patients enrolled into surgical trials).</i>  Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; and Final FIGO Stage {Cervical Cancer} coded as IB1, and Surgery Performed not coded as Trachelectomy, Radical Trachelectomy or Patient Refused Treatment; and Type of Systemic Anti-Cancer Therapy (SACT) 1 not coded as neo adjuvant; and Patient entered into Clinical Trial not coded as Yes – Surgical Trial.  [FIGO = IB1] <b>AND</b> [SURG <> 03 OR 04 OR 95] <b>AND</b> [CHEMTYPE1 <> 01] <b>AND</b> [TRIAL <> 01A]
Not recorded for numerator	Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target  [SURG = 99]
Not recorded for exclusion	Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.  [SURG = 99] <b>OR</b> [CHEMTYPE1 = 99] <b>OR</b> [TRIAL = 99]
Not recorded for denominator	Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard  [FIGO = 99]

## QPI 5 – Surgical Margins

<b>QPI Title:</b>	Patients with surgically treated cervical cancer should have clear resection margins.
<b>Description:</b>	Proportion of patients with cervical cancer who have surgical margins clear of tumour following hysterectomy.
Numerator	Number of patients with cervical cancer who undergo surgery where surgical margins are clear of tumour.  Margin Status {Cervical Cancer} coded as Clear  [MARGIN = 01]
Denominator	All patients with cervical cancer who undergo surgery. <i>(Excluding patients who undergo neo-adjuvant chemotherapy).</i>  Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; Surgery Performed coded as Hysterectomy or Radical Hysterectomy or Trachelectomy or Radical Trachelectomy; and Type of Systemic Anti-Cancer Therapy (SACT 1) not coded as Neoadjuvant.  ([SURG = 01 OR 02 OR 03 OR 04] <b>AND</b> [CHEMTYPE1 <> 01])
Not recorded for numerator	Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target  [MARGIN = 99]
Not recorded for exclusion	Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.  [CHEMTYPE1 = 99]
Not recorded for denominator	Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard  [SURG = 99]

## QPI 6 – 56 Day Treatment Time for Radical Radiotherapy

<b>QPI Title:</b>	Treatment time for patients with cervical cancer undergoing radical radiotherapy should be no more than 56 days.
<b>Description:</b>	Proportion of patients with cervical cancer undergoing radical radiotherapy whose overall treatment time, from the start to the end of treatment, is not more than 56 days.
Numerator	<p>Number of patients with cervical cancer undergoing radical radiotherapy (external beam or brachytherapy) whose overall treatment time, from start to end of treatment, is not more than 56 days.</p> <p>Date Treatment Completed (Radiotherapy) minus Date Treatment Started (Radiotherapy) coded as less than or equal to 56 days.</p> <p>( [RCOMPDATE – RSTARTDATE &lt;= 56])</p>
Denominator	<p>All patients with cervical cancer undergoing radical radiotherapy – (external beam or brachytherapy) (<i>No exclusions</i>).</p> <p>Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; and Radiotherapy Course Type coded as Radical or Chemoradiotherapy.</p> <p>[RADIOTYPE = 02 OR 06]</p>
Not recorded for numerator	<p>Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target</p> <p>[RSTARTDATE = 09/09/0909] <b>OR</b> [RCOMPDATE = 09/09/0909]</p>
Not recorded for exclusion	<p>Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.</p> <p>N/A</p>
Not recorded for denominator	<p>Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard</p> <p>[RADIOTYPE = 99]</p>



## QPI 7 – Chemoradiation

<b>QPI Title:</b>	Patients with cervical cancer undergoing radical radiotherapy should receive concurrent platinum-based chemotherapy.
<b>Description:</b>	Proportion of patients with cervical cancer undergoing radical radiotherapy who receive concurrent chemotherapy
Numerator	Number of patients with cervical cancer undergoing radical radiotherapy who receive concurrent chemotherapy.  Radiotherapy Course Type coded as Radical and Type of Systemic Anti-Cancer Therapy (SACT) 1-2 coded as Chemoradiotherapy.  [CHEMTYPE1 OR CHEMTYPE2 = 05]
Denominator	All patients with cervical cancer who undergo radical radiotherapy. <i>(No exclusions)</i> .  Date of Diagnosis [DIAGDATE] in range specified for comparative analysis; and Radiotherapy Course Type coded as Radical.  [RADIOTYPE = 02 OR 06]
Not recorded for numerator	Include in the denominator for measurement against the target. Present as not recorded only if the patient cannot otherwise be identified as having met/not met the target  [CHEMTYPE1 = 99] <b>OR</b> [RADIOTYPE = 99]
Not recorded for exclusion	Include in the denominator for measurement against the target unless there is other definitive evidence that the record should be excluded. Present as not recorded only where the record cannot otherwise be definitively identified as an inclusion/exclusion for this standard.  N/A
Not recorded for denominator	Exclude from the denominator for measurement against the target. Present as not recorded only where the patient cannot otherwise be definitively identified as an inclusion/exclusion for this standard  [RADIOTYPE = 99]