





# **FAQs**

# Catalogue of Requirements for Gynaecological Cancer Centres

of the German Cancer Society (Deutsche Krebsgesellschaft - DKG)

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Within the framework of the certification procedure, questions regularly crop up which require an explanation of the Technical and Medical Requirements. This document contains answers to the questions which the centres can refer to when implementing, and the experts can refer to when assessing the Technical and Medical Requirements.

#### Version FAQs and Catalogue of Requirements (CR)

Version status FAQ: 16 August 2024

The FAQs listed in this document are continuously checked to ensure that they are up to date and adapted in the event of changes to the Technical and Medical Requirements.

## **Overview of FAQs**

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## **Indicator Sheet**

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	Other carcinomas	
	Borderline Ovarian	
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## FAQs - Catalogue of Requirements Gyn

## 1.1 Structure of the network

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
1.1.3	<ul> <li>Gynaecological dysplasia units and consulting hours</li> <li>The separate certification of gynaecological dysplasia units and consulting hours can be done by the Gynaecological Cancer Centre or by one of its cooperation partners in line with the Catalogue of Requirements "Gynaecological Dysplasia". <a href="http://www.onko-zert.de/praxen_kooperationspartner.htm">http://www.onko-zert.de/praxen_kooperationspartner.htm</a></li> <li>Cooperation with certified gynaecological dysplasia units/consulting hours must be in place and the names must be given. Reasons for non-compliance are to be given separately.</li> </ul>	FAQ (27.08.2019) How is the requirement to be demonstrated?  Answer: If cooperation cannot be proven, the reasons must be explained in the audit. If the reasons are comprehensible (e.g. no certified dysplasia consultation/unit available within a radius of >45km or regionally related lack of incidence, etc.), there is no deviation.	

# 1.2 Interdisciplinary cooperation

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
1.2.1	Performance indicators Gynaecological Cancer	FAQ (14.07.2016)	
1.2.1	Centre	Is it correct that in the case of gynaecological tu- mours only the date of the postoperative histol-	
	Number of cases with a genital malignoma (i.e.	ogy "counts" as the initial diagnosis date, i.e. not	
	invasive neoplasias of the female genitals (no	the finding of the smear/pipelle de cornier/imag-	
	precancerous) borderline tumours of the ova-	ing procedures?	
	ries and serous tubal intraepithelial carcinoma	ing procedures.	
	(STIC)) per year:	Answer:	
	≥ 75 cases (= total case number), of which ≥	The counting date depends on the examination	
	50 primary cases	method that first gives the definitive diagnosis.	
		This can be a smear, but also the surgical histol-	
	Definition primary case:	ogy.	
	A primary case includes all stays and treat-		
	ments (surgery, radio(-chemo)therapy) of a		
	patient to treat a disease		
	Recurrence/metastasis of a patient is a new		
	case, not a primary case		
	Histology report, medical report and, where		
	appropriate, treatment/surgical report should		
	be available		
	<ul> <li>Planning/conduct of therapy via the Gynae-</li> </ul>		
	cological Cancer Centre Count time is the		
	time of the initial diagnosis or the time of the		
407	recurrence/metastasis	[5.0. (40.40.004 <del>7</del> )	
1.2.7	If a radiotherapy unit cooperates with several	FAQ (12.10.2017)	
	clinics, then all primary case patients with a	How should the requirement that all primary case	
	cervical carcinoma, who are to undergo radi-	patients with cervical carcinoma who are to be	
	ochemotherapy, should be presented in a cen-	treated with radiochemotherapy should present at	
	tre. To this end, the radiotherapy unit is to draw up a list of all patients presented to it that in-	one centre be interpreted?	
	cludes a centre assignment (certified centre,	Answer:	
	certification ongoing, not a centre). The	Patients who are primarily seen in radiation on-	
	presentation rate of 90% is to be achieved in	cology should be systematically brought to the tu-	
	each of the cooperating centres.	mour board. In order to facilitate the complete	
	odon or the ecoporating contros.	modification of the industrial complete	

## 1.2 Interdisciplinary cooperation

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
	This assignment of the patients is also of relevance for the tumour documentation.	presentation of these patients and their verifiability in the audit, a corresponding requirement was included in the data collection form (section	
		1.2.6.). The aim should be that the patients are presented in a certified Gynaecological Cancer Centre.	

## 1.4 Psycho-oncology

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre
1.4.2 b	Documentation and evaluation To identify treatment needs it is necessary to conduct standardised screening for mental strain (see Indicator "Psycho-oncological dis- tress screening"), and to document the result. The proportion of patients with excessive stress in the distress screening should be pre- sented.	FAQ (21.07.2016) Can an on-site contact replace screening?  Answer: No. In order to identify the need for treatment, it is necessary to carry out a <b>standardised</b> screening for psychological stress (see S3 guideline Psychooncology: e.g. Disress-Thermometer or HADS) and to document the result.
	Psycho-oncological counselling Psycho-oncological care, in particular for pa- tients with excessive stress in the distress screening, must be presented.	FAQ (28.08.2023) How should the proportion of patients with excessive distress in distress screening and further psycho-oncological care be presented?  Answer:
		The number of screened patients who have shown an excessive test should be described.
		The processes of psycho-oncological care should be described; the number of counselling sessions carried out should be recorded.
		See separate document FAQ Distress Screening.

## 1.6. Patient involvement

Section	Requirements	Explanatory remarks by the Cancer Centre
1.6.7	Event for patients	FAQ (26.05.2023)
	An information event for patients is to be staged by the Gynaecological Cancer Centre at least once a year. If patient events are (co-)financed by industry, this fact including potential conflicts of interest of the speakers must be disclosed. The centre must rule out any direct influence on pa-	How can the centre prove the exclusion of direct influence by industry representatives?  Answer:  Proof can be provided, for example, via internal compliance rules or, alternatively, via a self-decla-
	tients by industry representatives.	ration by the centre. In this, the centre should provide information on free access to the event, excluding the industry exhibition/information stands and remarks on contact between industry representatives and patrons.

## 1.7 Study management

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
1.7.5 c	Proportion study patients	FAQ (28.01.2022)	
1.7.5 0	1. Initial certification:	Do patients with gynaecological tumours who	
		were enrolled in the Heredi-CaRe study count to-	
	At the time of initial certification ≥ 1 patients must have been included in the studies.		
		wards the gynaecological cancer centre's student	
	2. After one year: at least 5% of the primary	quota?	
	case number	A	
	All study noticets can be taken into account	Answer:	
	All study patients can be taken into account	For the counting of HerediCaRe patients (proof of	
	when calculating the study rate (share study	study participation required), exclusive use of the	
	patients based on the Centre's primary case	checklist and referral of the patients to an FBREK	
	number).	centre is not sufficient.	
	Only the inclusion of patients in studies with an	FAO (40.00.0000)	
	ethical vote counts as study participation (non-	FAQ (10.02.2022)	
	interventional/diagnostic studies are also rec-	Can negatively screened study patients be	
	ognised).	counted?	
	General preconditions for the definition of the	A	
	study quota:	Answer:	
	<ul> <li>Patients can be counted once per study,</li> </ul>	Patients who have signed a informed consent	
	time: Date of patient consent	form for screening for study participation can be	
	All patients of the Centre can be counted	counted for the numerator of the respective study	
	<ul> <li>Study patients can be counted for 2 centres,</li> </ul>	indicator, even if the results of screening exami-	
	provided that the sending centre itself con-	nations performed with special diagnostics (no	
	ducts at least one study for patients of the	routine diagnostics) do not allow the patients to	
	Gynaecological Cancer Centre. If this	participate in the study.	
	method of counting is chosen (optional), the	EAO (05.07.0000)	
	centre must show how many patients are	FAQ (25.07.2022)	
	brought into studies at the centre itself, sent	Can studies with an ethical vote but without pa-	
	to other centres/clinics for study participation	tient informed consent - e.g. patient surveys - be	
	and taken over from other centres/clinics for	counted?	
	study participation.		
	<ul> <li>Registry studies can be counted if an ethics</li> </ul>	Answer:	
	vote and a study plan with a defined re-	No, these cannot be counted.	
	search question are available.	FAO (20 00 2022)	
	<ul> <li>Prevention/screening studies of the own</li> </ul>	FAQ (28.08.2023)	
	dysplasia consultation/unit can be counted	Can patients referred to a Centre for Personal-	
	for the own Gynaecological Cancer Centre	ised Medicine (CPM) for the purpose of complex	
		diagnostics, interdisciplinary consultation and in-	
		dividual therapy recommendations who partici-	
		pate in a study there be counted towards the	
		study quota of the sending centre?	
		Anavar	
		Answer:	
		Yes, in this case the study inclusion can be	
		counted by both the sending centre and the	
		CPM. The other requirements for study inclusion	
		according to the survey form will apply.	

## 2.1 Consulting hours

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
2.1.7	Hereditary stress	FAQ (26.05.2023)	
	Cooperation with certified centres for familial breast and ovarian cancer (FBREK centres) for counselling and genetic testing must be demonstrated in writing in accordance with the	Does the non-fulfilment of the requirement "Co- operation with certified centres for familial breast	

# 2.1 Consulting hours

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
	FBREK (familial breast and ovarian cancer) co-	and ovarian cancer (FREBK centres) for counsel-	
	operation agreement of the vdek (=Association	ling and genetic testing must be demonstrated."	
	of substitute health insurance funds)	result in a deviation?	
	Check lists to record hereditary stress are to be applied in the case of:  Patients with breast/ovarian cancer (mainly familial breast/ovarian cancer)  Patients with endometrial cancer (EC) (mainly HNPCC/Lynch syndrome	Answer: A written cooperation with an FBREK centre must be bindingly proven.	
	The current check lists and the algorithm can be downloaded from this <u>Link</u> in the section Gynaecological types of cancer.		

## 5.2 Organ-specific surgical therapy

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre
Section 5.2.1	Requirements  Specialists for the Gynaecological Cancer Centre  • At least 2 specialists for gynaecology with the focus designation Gynaecological Oncology in line with the staffing schedule working for the Gynaecological Cancer Centre  • The names of the specialists are to be given.  Initial certification: At least 1 specialist for gynaecology with the focus designation Gynaecological Oncology. A second specialist for gynaecology should be undergoing specialty training for the focus designation Gynaecological Oncology. This must have been successfully concluded before recertification (after 3 years) and notified.  A concept for the training of gynaecological on- cology specialists must be available. In addi- tion, the doctors undergoing training (+ proof of logbook) should be named. Deviations should be justified.	Explanatory remarks of the Gyn. Cancer Centre  FAQ (14.07.2016)  What is the procedure to be followed in the event of the departure/absence of the second focal point holder?  Answer:  If no second focal point holder is available for the centre after re-certification (e.g. departure/absence), the replacement must take place within 12 months of the date of departure/absence.  FAQ (14.07.2016)  What should be done if there is no evidence of a second focal point holder at the time of recertification?  Answer:  It must be proven that activities to establish a second focal point holder took place after the initial certification (e.g. new appointment, training,).  The reasons for the lack of a second focal point
	focus designation Gynaecological Oncology. A second specialist for gynaecology should be undergoing specialty training for the focus designation Gynaecological Oncology. This must have been successfully concluded before recertification (after 3 years) and notified.  A concept for the training of gynaecological oncology specialists must be available. In addition, the doctors undergoing training (+ proof of logbook) should be named. Deviations should	What should be done if there is no evidence of a second focal point holder at the time of recertification?  Answer: It must be proven that activities to establish a second focal point holder took place after the initial certification (e.g. new appointment, training,).

## 5.2 Organ-specific surgical therapy

Requirements	Explanatory remarks of the Gyn. Cancer Centre
requirements	staffing plan in relation to their work in the Gynaecological Cancer Centre. How is the specification "according to the staffing plan in activity for the Gynaecological Cancer Centre" to be understood? What scope of activity is to be demonstrated?
	Answer: This formulation means that both specialists must regularly work for the Gynaecological Cancer Centre, which also takes into account deputising arrangements (guideline: 0.5 HC/specialist with a focus on the Gynaecological Centre). A substitute on an hourly basis is not sufficient. For a positive evaluation, a concrete description of the activities of the specialist with a specialisation is required (detailed naming in the questionnaire). At the time of re-certification, the involvement of the second specialist with a speciality must be proven for at least three months.
	FAQ (26.05.2023) Can the "Optional further training in special surgical gynaecology" according to the (model) further training regulations (MWBO) 1992 be recognised in terms of the specialisation in gynaecological oncology?
	Answer: Yes, it can be recognised if it is recognised as equivalent by the State Medical Association. There must be at least 1 specialist in Gynaecology with a specialisation in Gynaecology.
20 surgeries a year, also possible when senior surgeon supervises surgery as an assisting	FAQ (26.05.2023) How are the surgeries to be counted for the surgeons?
All surgical cases of the GC must be operated on by designated surgeons (first surgeon or as training assistant).	Answer: All surgeries that are counted for the implementation of indicator 7 (operated cases with genital malignancy) can be assigned to 1 surgeon. Any difference between the sum of "Operations per named surgeon" and the "Operative cases" indicator must be explained (e.g. surplus from the year before the indicator year).
	FAQ (15.05.2019) Who can be appointed as an operator?
	Answer: A gynaecology specialist who fulfils the quantitative requirements (at least 20 operations per year) and is at least in further training to become a specialist (proof of expertise: certificate from the head of the centre).
	Number of surgeries per named operator: 20 surgeries a year, also possible when senior surgeon supervises surgery as an assisting surgeon.  All surgical cases of the GC must be operated on by designated surgeons (first surgeon or as

## 5.2 Organ-specific surgical therapy

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre
		FAQ (26.05.2023)
		Is there a specification as to which surgical pro-
		cedures can be counted? - E.g. laparoscopy with
		PEs to secure an advanced ovarian carcinoma
		(1) - lymph node staging for cervical carcinoma
		as a surgical case if followed by radiotherapy (2)
		- HSK/ curettage for endometrial cancer (3)?
		Answer:
		For 1) and 2): Counts as a surgical case, To 3):
		Does not count if only on the basis of diagnosis.

## 6.2 Organ-specific medicinal oncological therapy

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre	
6.2.3	Qualification treatment unit/partner	FAQ (12.10.2017)	
	at least 50 drug-based tumour therapies	Can patients who receive both chemotherapy	
	(cytostatic therapies and / or targeted ther-	and antibody therapy be counted twice for the	
	apeutics and / or antibody / immune thera-	treatment unit expertise?	
	pies, no hormone therapies) every year in		
	the case of patients with gynaecological /	Answer:	
	senologic forms of cancer	If chemotherapy and AK therapy are adminis-	
	or	tered in parallel, the patient cannot be counted	
	at least 200 drug-based tumour therapies (cytostatic therapies and / or targeted therapeutics and / or antibody / immune therapies, no hormone therapies) every year (in the case of different types of tumour)	twice.	
	Calculation method: completed systematic/ cytostatic / targeted therapy per patient (consisting of several cycles or administrations).		
	When this number is not reached, expertise cannot be proven by means of cooperation.		

## 8 Pathology

Section	Requirements	Explanatory remarks of the Gyn. Cancer Centre
8.4	Specialists - Expertise	FAQ (17.08.2021)
	20 histologies/year per designated specialist (incl. PE)	What histologies can be counted?
		Response:
		Only histologies of invasive neo-plasias of the fe-
		male genitalia, borderline tu-mors of the ovary
		(BOT) and serous tubular intraepithelial carcino-
		mas (STIC) can be counted, not histologies of
		precancerous lesions <u>.</u>

#### FAQs - Indicator Sheet Gyn

#### **Basic Data Columns A-I:**

#### FAQ (14.07.2016)

Do dysgerminomas of the ovary and sarcomas count as other carcinomas?

Answer: Yes.

#### FAQ (14.07.2016)

What counts as non-cancerous ovaries?

Answer: Germ cell tumours and germ cell stromal tumours.

#### FAQ (12.10.2017)

Does carcinosarcoma of the ovary count as ovarian carcino-men or as other tumours?

Answer:Other tumours.

#### FAQ (12.10.2017)

Does a malignant melanoma of the vulva count as a primary case for the Gyn. Cancer Centre?

Answer:

No, it cannot be counted.

#### FAQ (21.08.2018)

Does basal cell carcinoma of the vulva count as a vulvar carcinoma?

Answer:

Yes, it counts as a vulvar carcinoma. Only for code 26 (inguinofemoral staging) it is not counted according to the definition of the code.

### FAQ (21.08.2018)

Do **dermoid cysts of the ovary** (ICD-O-M 9084/0) count as primary cases for the Gyn. Cancer Centre? Answer

No, these cannot be counted.

#### FAQ (27.08.2019)

Does malignant mixed müllerian tumour count as other carcinoma?

Answer:

Yes.

### FAQ (12.10.2017)

Do **borderline tumours of the ovary** also include those with the dignity "uncertain behaviour (ICD-10 D39.1)? Answer:Yes, these are counted as BOT.

### FAQ (12.10.2017)

Do operated patients with ovarian cancer without R0 resection have to be shown in column D "**Not complete surgery**"?

Answer:No. Patients with definitive surgery and R1 resection are to be shown in column E "Definitive surgery = staging surgery". In column D "Incomplete surgery", only those patients are shown who prove to be inoperable during the surgical intervention.

### FAQ (14.11.2017)

Can primary **peritoneal carcinomas** (ICD-10 C48) be counted as primary cases? Answer :Yes.

### FAQ (27.08.2019)

Is it sufficient if the **recurrence of an ovarian carcinoma** is diagnosed solely on the basis of a resurgent tumour marker and imaging suspicion of a recurrence, or is histological confirmation always required as well? Answer:In the case of ovarian carcinoma, imaging and/or tumour markers are sufficient; histological confirmation is not obligatory.



#### FAQ (29.06.2020)

Can patients with **SEIC** (serous endometrioid in-traepithelial carcinoma) be counted for the Gyn. Cancer Centre be counted?

Answer: Yes, they can be counted.

#### FAQ (02.07.2020)

Can extramammary Paget's disease of the vulva be counted as a primary case?

Answer: No, it cannot be counted.

#### FAQ (17.08.2021)

Does a goiter carcinoid of the ovary (morphology code: 9091/1) count as "other cases"?

Answer: No, it does not count because it is benign.

#### FAQ (17.08.2021)

Does a granulosa cell tumour of the ovaries count as a primary case?

Answer: A granulosa cell tumour with ICD-O-M 8620/1 does not count, only the malignant granulosa cell tumour with ICD-O-M 8620/3. The latter counts as "other cases".

### FAQ (17.08.2021)

How should a **bilateral mucinous ovarian carcinoma**, one with a **proportion of borderline tumour**, be documented in the indicator sheet?

Answer: The patient is evaluated as one primary case despite the fact that she has both tumours. The FIGO stage of the mucinous ovarian carcinoma and not the borderline tumour is decisive for the entry in the data sheet.

#### FAQ (17.08.2021)

Does an angiomyxoma of the vulva count as a primary case?

Answer:

No, only inv. Neoplasms of the female genital tract (incl. BOT and STIC) can be counted.

#### FAQ (03.05.2023)

Does a malignant GIST count for the Gynaecological Cancer Centre?

Answer: Yes, it counts as an "Other case".

### FAQ (03.05.2023):

Does an **epithelioid sarcoma / myoepithelial differentiated tumour of the mons pubis** count for the Gynae-cological Cancer Centre?

Answer: Yes, it counts as an "Other case".

#### FAQ (10.05.2023):

Can **primary peritoneal mesotheliomas** be counted as primary cases for the Gynaecological Cancer Centre? Answer: No. they cannot be counted. See also: Mesothelioma units certification system.

#### FAQ (10.05.2023)

Can STIL (serous tubular intraepithelial lesion) of the ovary be counted in addition to STIC?

Answer: No.

#### FAQ (10.05.2023)

Does large **cell neuroendocrine (LCNEC) corpus carcinoma** count for the Gynaecological Cancer Centre? Answer: Yes, it counts as an "Other case".

#### FAQ (10.05.2023)

Does a **neuroendocrine cancer of the ovary** (large cell neuroendocrine cancer, 8013/3) count for the Gynae-cological Cancer Centre?

Answer: Yes, it counts as an "Other case".

### FAQ (10.05.2023)

How do patients with cervical carcinoma who undergo brachytherapy at the Gynaecological Cancer Centre count? Cancer Centre have received brachytherapy only?

Answer: Patients who only receive brachytherapy at the centre and no other measures such as TC cannot be counted as (primary) cases for the centre.

## FAQ (16.05.2023)

Do germinally mixed tumours (9085/3) of the ovary count as cases for the Gynaecological Cancer Centre?

Answer: Yes, they count as "Other cases".

## **Basic Data Columns J-K:**

#### FAQ (24.05.2016)

Can non-primary cases also include **progressions**?

#### Answer:

No, progressions cannot be counted.

	T -	1	1 -	T
9	Surgical staging early ovarian cancer	Numerator	Primary cases of the denominator with surgical staging with: •Laparotomy •Peritoneal cytology •Peritoneal biopsies •Bilateral adnex exstirpation •Hysterectomy, where appropriate extraperitoneales procedure •Omentectomy at least infracolic •Bilateral pelvic and paraaortal lymphonodectomy	FAQ (14.07.2016) Peritoneal biopsies should be performed even if the peritoneum is macroscopically unremarkable. Macroscopically unremarkable peritoneum is not sufficient justification for not performing biopsies. In these cases, a deviation should be pronounced.  FAQ (10.05.2023) Does surgical staging have to be performed in one session or can it also be performed in
		Denominator	Surgical primary cases ovarian cancer FIGO I – IIIA	two sessions?  Answer:
		Target value	No target value	Both a one-session and a two- session procedure are permit- ted.
10	Macroscopic complete resection advanced ovarian cancer	Numerator	Primary cases of the de- nominator with macroscopic complete resection	FAQ (25.07.2016) What does "macroscopically complete resection" mean?
		Denominator	Surgical primary cases with an ovarian cancer FIGO IIB-IV	Answer: The final ope-rative result is < R2, i.e. R0 or R1.
		Target value	≥ 30%	FAQ (14.07.2016) In the case of multiple operations, does the macroscopically complete resection refer to the first tumour-specific operation or also to the last tumour-specific operation on the tumour?
				Answer: The macroscopically complete resection is decisive, regardless of the number of operations



11	Operation advanced	Numerator	Primary cases of the de-	FAQ (14.07.2016)
	ovarian cancer by a gynaecological oncol- ogist	Numerator	nominator whose definitive surgical treatment was performed by a gynaecological oncologist	The operations were performed by a gynaecological oncologist as a training assistant. The main surgeon was
		Denominator	Surgical primary cases ovarian cancer FIGO IIB-IV after completion of surgial treatment	not a gynaecological oncologist. Can the operations still be included in the numerator?
		Target value	≥ 80%	Answer: Yes.
13	First-line chemother- apy advanced ovarian cancer	Numerator	Primary cases of the de- nominator with first-line chemotherapy with car- boplatin and paclitaxel	FAQ (10.05.2023) Can patients who receive additional substances - e.g. as part of a study - be counted in the
		Denominator	Primary cases ovarian can- cer FIGO IIA-IV	numerator?
		Target value	No target value	Answer: Yes, these can be counted.
				FAQ (10.05.2023)  Does the administration of carboplatin/paclitaxcel refer to adjuvant or neoadjuvant administration?
				Answer: to the adjuvant administration.
14	Cytological/ histological lymph node staging	Numerator	Primary cases of the de- nominator with cytologi- cal/histological lymph node staging	FAQ (14.07.2016) The numerator can include both primary cases with cytologic/histologic lymph node
		Denominator	Primary cases cervical can- cer FIGO stage ≥ IA2-IVA	staging as part of diagnosis and primary cases with thera-
		Target value	≥ 60%	peutic lymph node removal as part of surgical treatment. LN staging in the context of diagnostics as well as primary cases with therapeutic lymph node removal in the context of surgical therapy can be taken into account.
				FAQ (12.10.2017) Can pure imaging LN staging be counted for the indicator?
				Answer: No, this does not count for the indicator.
15	Brachytherapy as a component of primary radio(chemo) therapy	Numerator	Primary cases of the de- nominator in which brachy- therapy was administered as part of primary ra- dio(chemo) therapy	FAQ (17.08.2021) What is meant by primary radio(chemo)therapy? Answer:
		Denominator	Primary cases with cervical cancer and primary radio(chemo) therapy, without primary Distant Metastasis	The intention of primary radio(chemo)therapy (= radiochth planned as the first and only therapy) is decisive for

		Target value	≥ 80%	the denominator. In exceptional cases, a so-called secondary (not primarily planned) hysterectomy or so-called extended chemotherapy may be performed, but ultimately this is irrelevant for the denominator, because these patients can also be counted.  FAQ (17.08.2021) Can brachytherapy equivalents such as Cyberknife or Boost also be counted?  Answer:
				No, they cannot be counted.
17	Details in pathology report in the case of first diagnosis and tu- mour resection	Numerator	Primary cases of the de- nominator with pathology reports containing details of:	FAQ (12.10.2017) Does the pTNM (staging) have to be complete?
			Histological type according to WHO,     Grading,     Detection/non-detection of lymph or blood vessel infiltration (L and V status),     Detection/non-detection of perineural invasion (Pn status)	Answer: The key figure refers to the content of the pathological report. If no lymph node removal was performed, no pN can be given. cN cannot be a substitute because it was not determined by the pathologist.
			tus), •Staging (pTNM), •Depth of invasion and spread in mm in the case of pT1a, three-dimensional tu- mour size in cm (ab pT1b), •Metric details of the mini-	FAQ (12.10.2017) Is a separate resection margin expected here from the invasive carcinoma and VIN respectively?
			mum distance of the carcinoma and VIN from the vulvar resection margin in the histological specimen; •In the case of resection of	Answer: Yes, separate indication of the resection margin of invasive carcinoma and VIN.
			the vulvar-vaginal or vulvar- anal transition zone and, where applicable, of the urethra metric details of the minimum distance to the	FAQ (12.10.2017) Are VIN III lesions to be considered or are VIN I and VIN II also included?
			vulvar-vaginal or vulvar- anal and, where applicable, urethral resection margin; •Metric details of the mini- mum distance to the soft	Answer: The guideline only specifies VIN, therefore VIN I-III are meant.
			tissue resection margin (ba-	FAQ (17.08.2021)
		Denominator	sal margin) Primary cases vulvar cancer with tumour resection	How is the three-dimensional tumour size to be indicated?
		Target value	≥ 80%	Answer: Three-dimensional tumour size in cm = length in cm (horizonal extension) x width in cm (vertical extension) x depth in cm

				(infiltration depth). But it is not the cubic centimetres that are asked for, but the extent of the expansion, i.e. cm in each case.
18	Details in pathology report in the case of lymphonodectomy	Numerator	Primary cases of the denominator with pathology report with details of:  • Number of affected lymph nodes in relation to the number of removed lymph nodes classified by removal localisation (inguinal/pelvic)  • Non-detection/detection of a capsel infiltration of the lymph node metastatis and/or detection lymph node infiltrations in perinodal fatty tissue and/or the lymph node capsule (>=pN2c)  • Biggest spread of metastases (through pN details)	FAQ (27.08.2019) Are patients with only sentinel lymphonodectomy (without conventional LNE) taken into account here?  Answer: No.
		Denominator  Target value	Primary cases vulvar cancer with lymphonodectomy ≥ 80%	
19	Conduct inguinofemoral staging	Numerator	Primary cases of the de- nominator with surgical staging (systematic lym- phadenectomy and sentinel biobsy) of inguinofemoral lymph nodes	FAQ (12.10.2017) Which operation codes are to be documented for this key figure? Answer:
		Denominator	Primary cases vulvar cancer ≥ pT1b (no basal cell carcinoma and no verrucous carcinoma)	It concerns lymph node staging, which is usually coded with its own OPS. There are several OPS that
		Target value	≥ 90%	can be used for this, depending on the operation performed. The surgeons are responsible for entering these OPSs, if necessary in consultation with Controlling.
20	Sentinel lymph nodes biopsy	Numerator	Primary cases of the denominator with the following characteristics:  • Clinical tumour size < 4 cm and  • Unifocal tumour (= no multiple tumours; TNM m-symbol) and  • Clinically inconspicuous lymph nodes (cN0) and  • Pathohistological ultrastaging of lymph nodes (= in line with LL), only if all sentinel lymph nodes are tumor-free in the H&E staining	FAQ (17.08.2021) What is pathohistological ultrastaging?  Answer: Ultrastaging, i.e. the immunohistochemical examination of the lymph nodes with a pancytokeratin antibody, is carried out if all sentinel lymph nodes are negative in the HE stain. If the LK are positive in the conventional staining (= HE), no ultrastaging is carried out.



	T	Denominator	Primary cases vulvar can	
		Denominator	Primary cases vulvar can- cer and sentinel lymph	
			node biopsy	
		Target value	≥ 80%	
		Target value	2 00 70	
26a	Hysterectomy without	Numerator	Primary cases of the de-	
26b	morcellement for sar-		nominator with hysterec-	
	coma confined to the		tomy without morcellement	
	uterus	Denominator	25a: Cases operated on at	FAQ (19.11.2021)
	(25a: in the centre,		the centre Primary cases	Which morphology codes
	25b: outside of the		with sarcoma confined to	count?
	centre)		the uterus (ICD-O T C54,	
			C55 iVm morphology codes	Answer
			sarcoma centres), M0 with	Morphology codes 8930/3
			hysterectomy	(high grade endometrial stro- mal sarcoma) and 8931/3 (low
			25b: Primary cases oper-	grade endometrial stromal sar-
			ated on outside the centre	coma) count.
			with sarcoma confined to	33
			the uterus (ICD-O T C54,	
			C55 iVm morphology codes	FAQ (26.04.2022)
			sarcoma centres), M0 with	What does "primary cases op-
			hysterectomy	erated on outside the centre"
		Target value	No target value	mean?
				Answer
				This means, for example, pa-
				tients who have had a hyster- ectomy outside the centre,
				who have evidence of a sar-
				coma in the histology and who
				then come to the centre and
				are primary cases of the cen-
				tre because the centre takes
				over the therapy and further
				care of the patients.
				FAQ (26.05.2023)
				How is the numerator of the in-
				dicators to be understood?
				A
				Answer:
				Only sarcomas that were re-
				moved by hysterectomy WITHOUT morcellement are
				counted for the indicators. In-
				side (=a) or outside (=b) the
				GC.
		1		