

# 子宮頸癌診療指引

## 婦癌多專科團隊

2005年05月制定 2010年08月修訂

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2013年08月修訂 2014年12月修訂

2015年12月修訂 2016年07月修訂

參考資料：

Cervical Cancer Guidelines V2.2015

全民健康保險藥品給付規定行政院衛生署一零五年版(30051\_2)

Physicians' Cancer Chemotherapy Drug Manual 2010

LCIS = Lobular carcinoma in situ

DCIS = Ductal carcinoma in situ

(+) = positive

(-) = Negative

LN = lymph node

R/T = radiation therapy

$\overline{c}$  With

$\overline{s}$  = without

ALP= alkaline phosphatase

PBI = partial breast irradiation

CR =Complete response

PD =Progressive disease

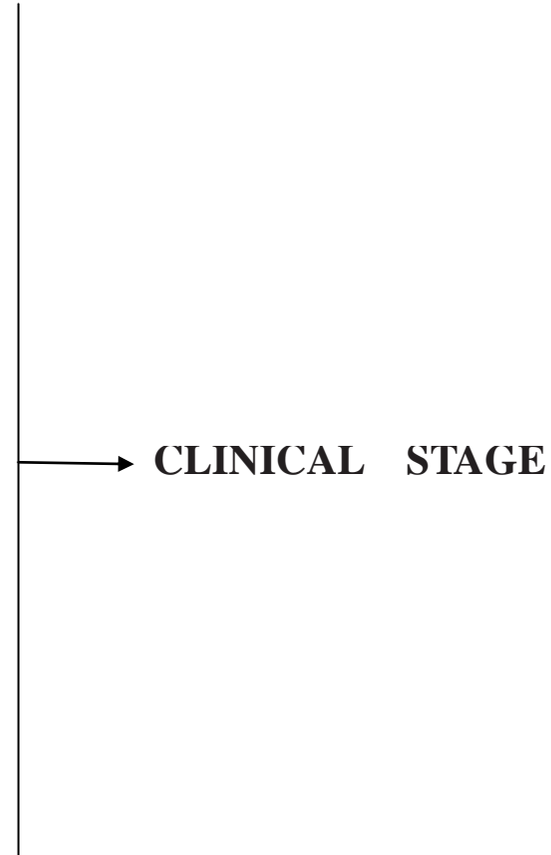
LVSI =Lymphovascular space invasion

## WORK UP

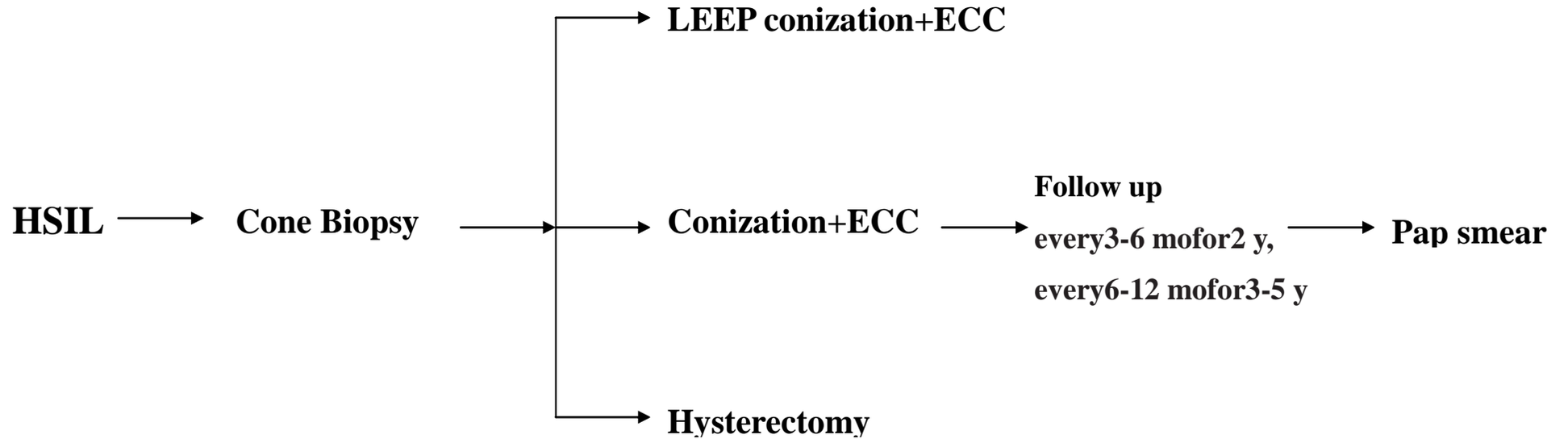
- History
- Physical examination
- Complete blood count(CBC) & Platelet
- Pathologic Review
  - Cervical biopsy or cone biopsy
- Liver function test/Renal function studies
- Imaging

(Optional for  $\leq$  stage IB1):

  - Chest x-ray
  - CT
  - MRI as indicated

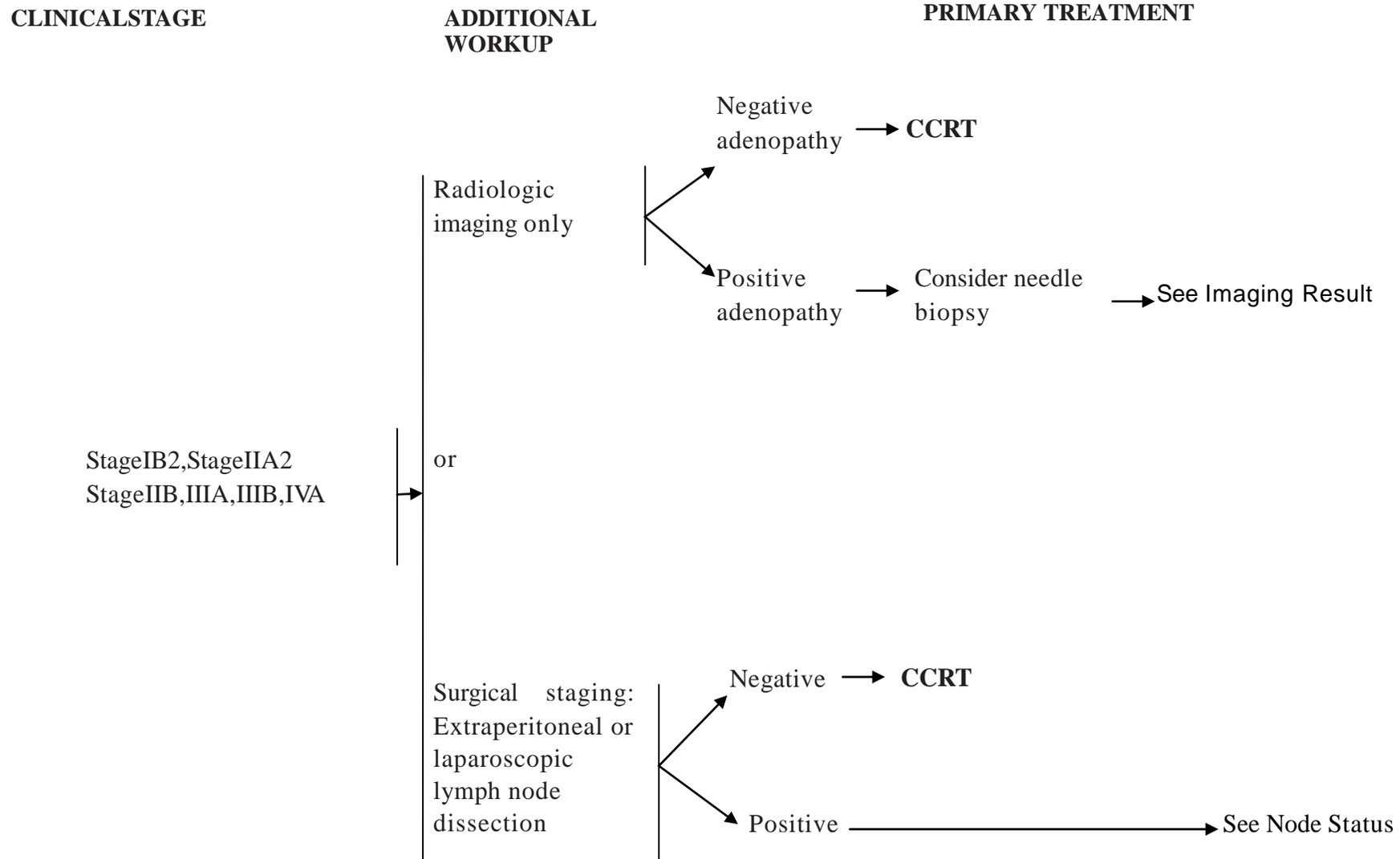


## TREATMENT OF HSIL (CIN III)



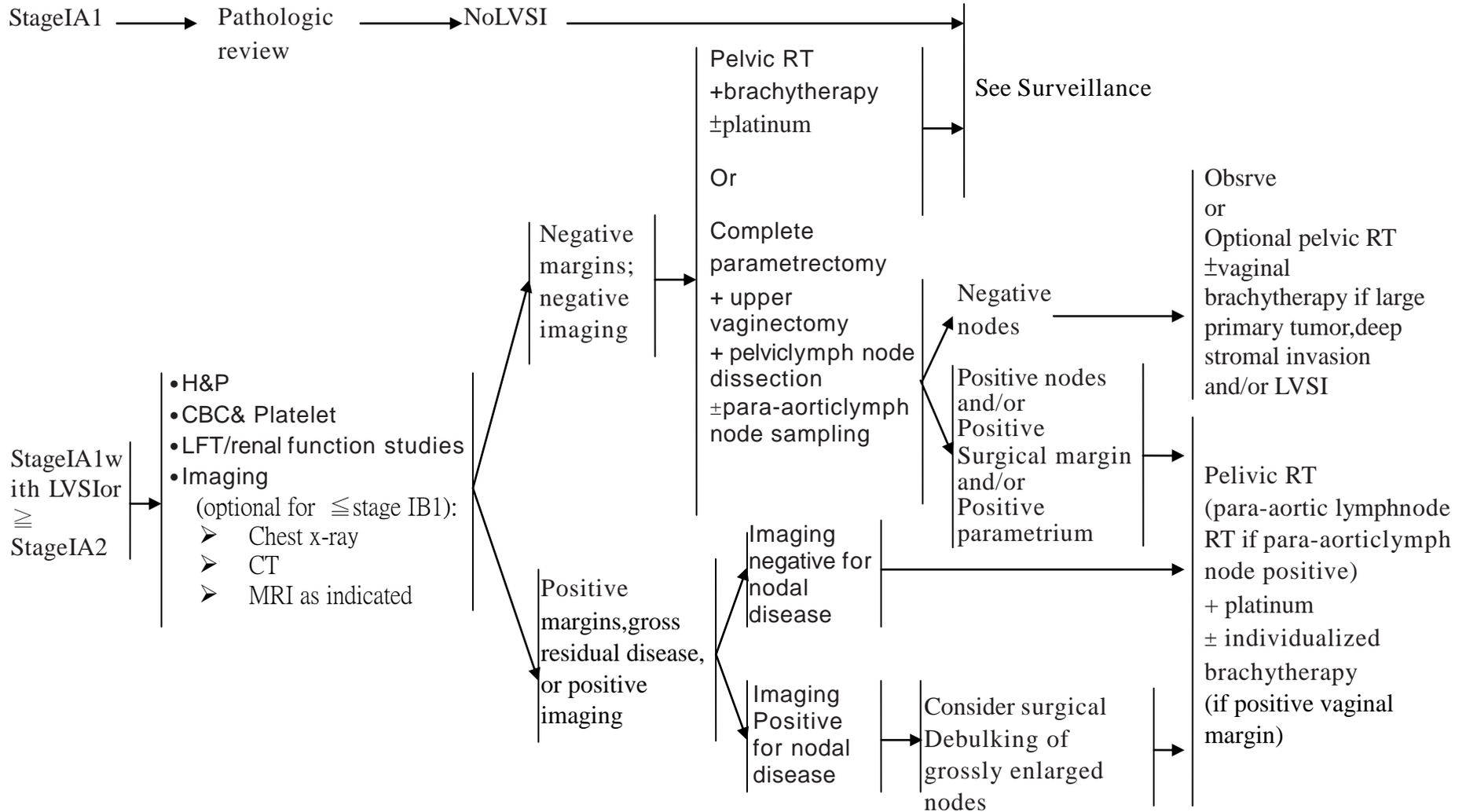
CLINICAL STAGE		PRIMARY TREATMENT	
Stage IA1 (No LVSI)	Fertility sparing	<b><u>Cone biopsy with negative margins:</u></b> (preferably a non-fragmented specimen with 3-mm negative margins) (If positive margins, repeat cone biopsy or perform trachelectomy)	Surveillance
	Non-Fertility sparing	<b><u>Cone biopsy with Negative margins and inoperable:</u></b> Observe	
		<b><u>Cone biopsy with Negative margin sandoperable:</u></b> Extrafascial hysterectomy	
Stage IA1 (LVSI) and Stage IA2	Fertility sparing	<b><u>Cone biopsy with Positive margins for dysplasia for carcinoma:</u></b> modified radical hysterectomy+ pelvic lymph node dissection	Surgical Findings
		1. Negative margins Cone biopsy is enough.	
	2. If positive margins, repeat cone biopsy or perform trachelectomy+ pelvic lymph node dissection ± para-aortic lymph node sampling (2B)	Surveillance	
Non-Fertility sparing	Pelvic RT+brachytherapy (total point A dose 70~80Gy)	Modified radical hysterectomy + pelvic lymph node dissection	Surgical Findings

CLINICAL STAGE		PRIMARY TREATMENT	
Stage IB1	Fertility sparing	Radical trachelectomy + pelvic lymph node dissection	Surveillance
	Non-Fertility sparing	Radical hysterectomy + pelvic lymph node dissection	Surgical Findings
Stage IIA1	Non-Fertility sparing	Or Pelvic RT +brachytherapy(total point A dose 80~85Gy) ±concurrent CCRT	Surveillance
Stage IB2 and Stage IIA2	Non-Fertility sparing	Definitive Pelvic RT 1.8-2G perfraction (4-6cycles) + ciplatin 40mg/ms2/week +brachytherapy(total point A dose $\geq$ 85GY)	Surveillance
		Radical hysterectomy + pelvic lymph node dissection	Surgical Findings
		Pelvic RT + platinum +brachytherapy(total point A dose 75-80GY) +adjuvant hysterectomy	Surveillance



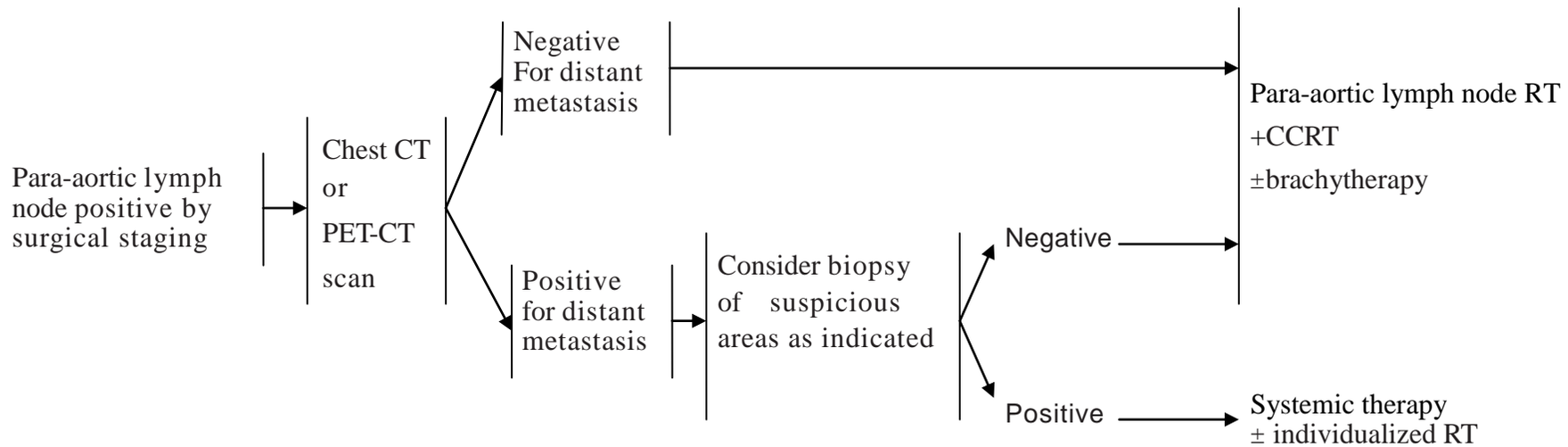
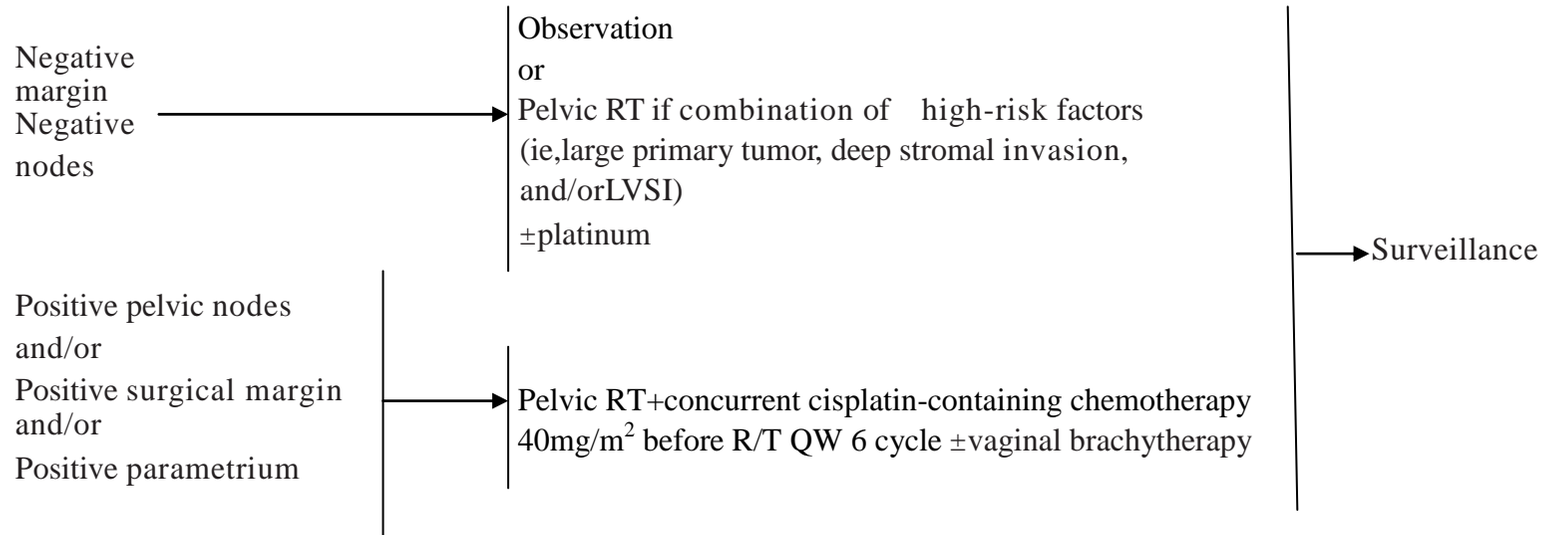
INCIDENTAL FINDING OF INVASIVE CANCER AT SIMPLE HYSTERECTOMY

PRIMARY TREATMENT



**SURGICAL FINDINGS**

**ADJUVANT TREATMENT**

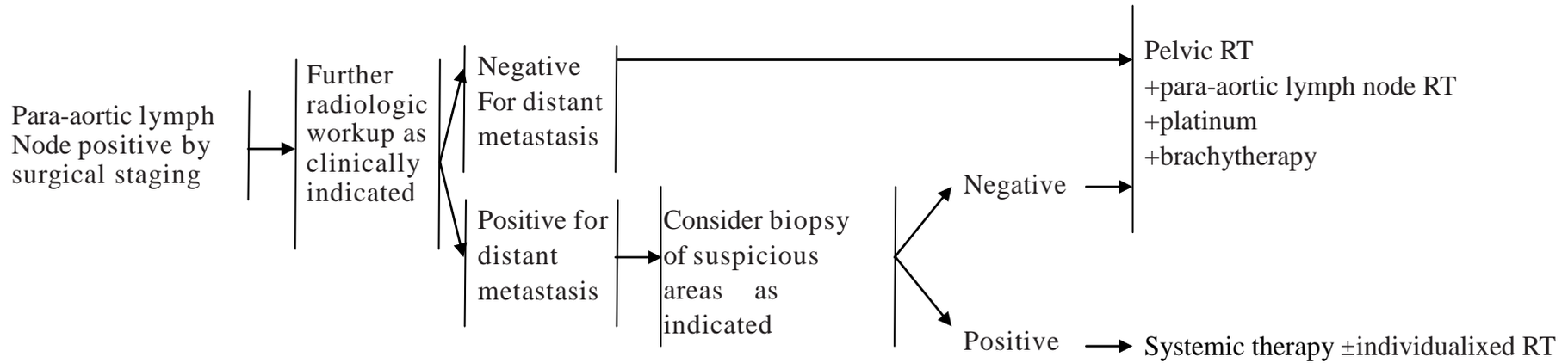


Stage IB2, IIA2; Stage IIB, IIIA, IIIB, IVA  
 NODE STATUS

PRIMARY TREATMENT

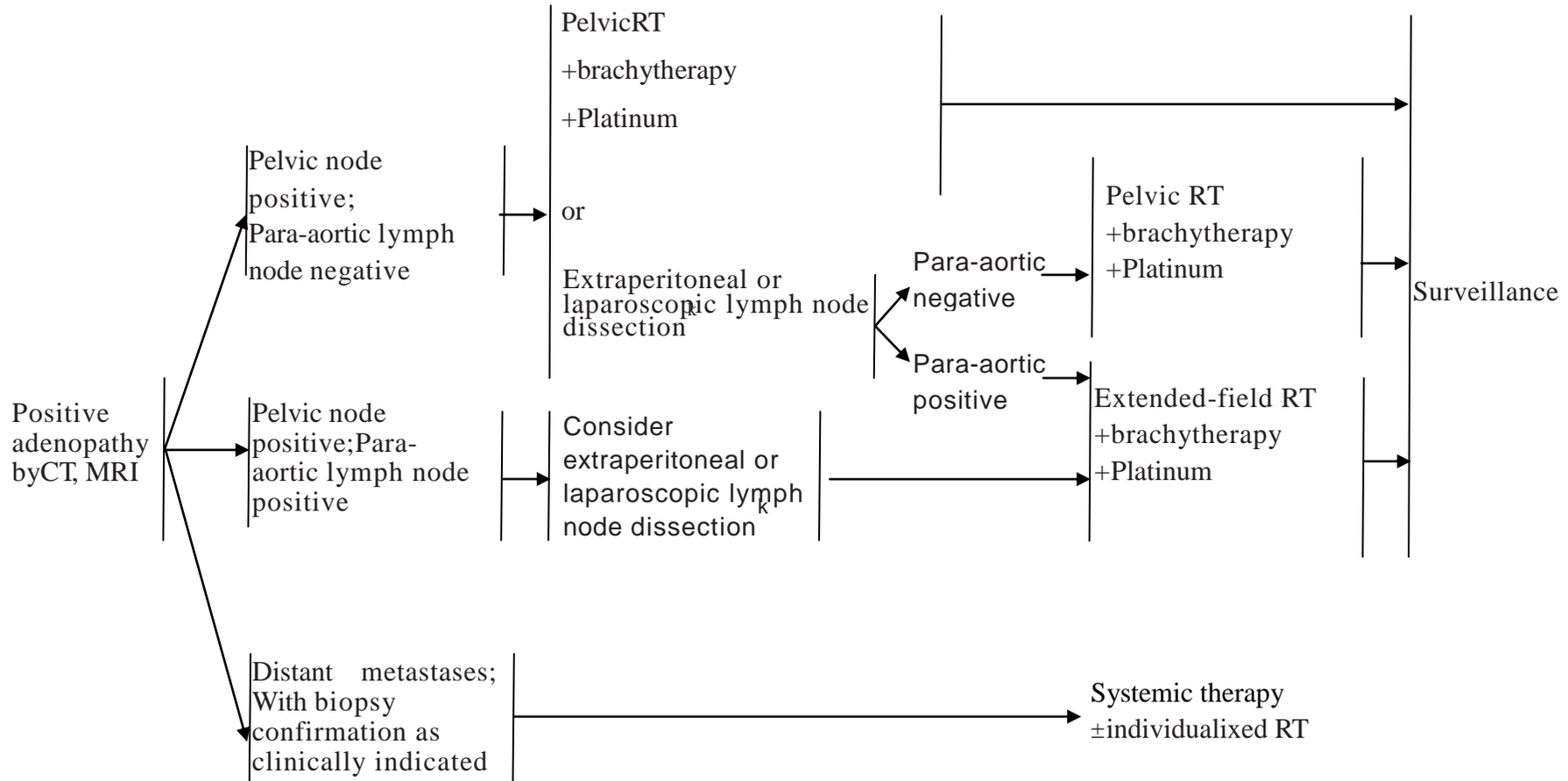
Pelvic lymph node positive  
 and para-aortic lymph  
 node negative by surgical  
 staging

CCRT  
 +brachytherapy



Stage IB2, IIA2  
 Stage IIB, IIIA, IIIB, IVA  
 IMAGING RESULTS

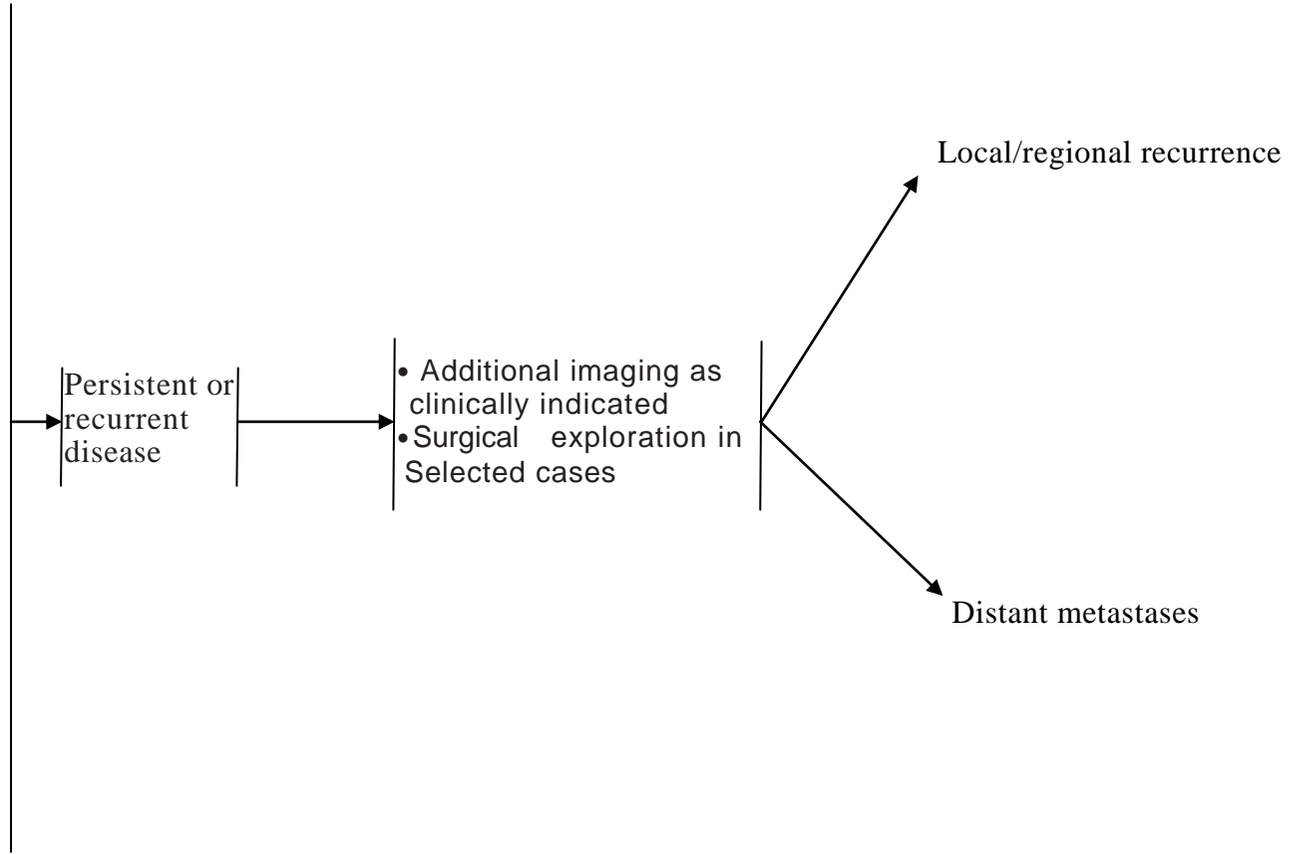
PRIMARY TREATMENT



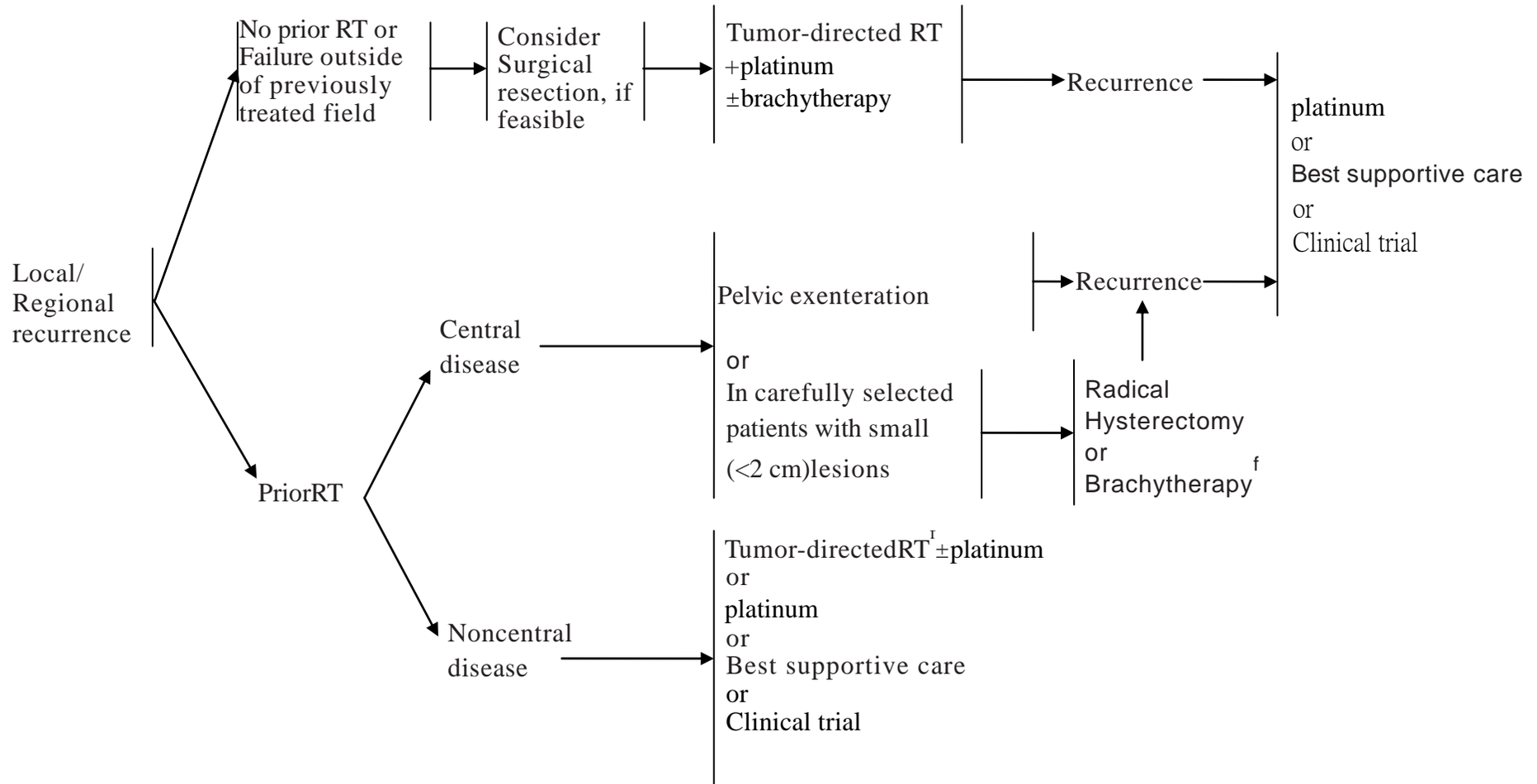
SURVEILLANCE

WORKUP

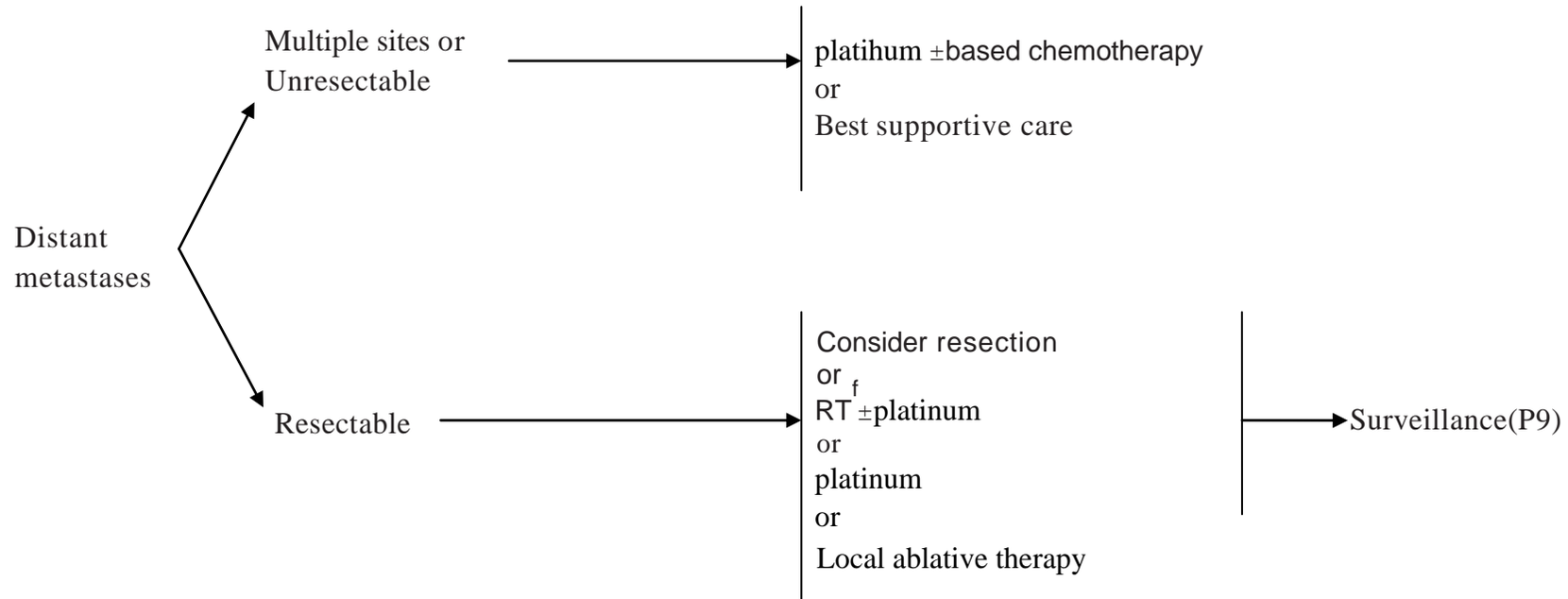
- Interval History & Physical
- Cervical/vaginalcytology  
Examination every 3-6 mo for 2 y,  
then every 6-12 mo for 3-5 y,  
then annually (based on patient's risk of  
disease recurrence).
- Imaging
  - Chest radiography
  - CT, MRI as clinically indicated for  
suspicious recurrence.
- Laboratory assessment
  - CBC & Platelet
  - BUN/creatinine findings suspicious  
for recurrence
- Recommend use of vaginal dilator  
after RT
- sexual health
- Patient education regarding symptoms,  
life style
- Obesity, exercise



THERAPY FOR RELAPSE



THERAPY FOR RELAPSE



一、化學治療

**CHEMOTHERAPY REGIMENS FOR RECURRENT OR METASTATIC CERVICAL CANCER<sup>†</sup>**  
(Strongly consider clinical trial)

**First-line combination therapy**

- Cisplatin/paclitaxel+bevacizumab<sup>1</sup>
- Carboplatin/paclitaxel<sup>4,5</sup>
- Cisplatin/topotecan+bevacizumab<sup>6</sup>
- Cisplatin/gemcitabine(category3)<sup>7</sup>
- Cisplatin/paclitaxel(category1)<sup>2,3</sup>

**Possible first-line single-agent therapy**

- Cisplatin (preferred as a single agent)<sup>3</sup>
- Carboplatin<sup>8</sup>
- Paclitaxel<sup>9</sup>

**Second-line therapy**

(Agents listed are category 2B unless otherwise noted)

- Bevacizumab
- Docetaxel
- 5-FU (5-fluorouracil)
- Gemcitabine
- Ifosfamide
- Irinotecan
- Mitomycin
- Topotecan
- Pemetrexed(category 3)
- Vinorelbine(category 3)

1. Bevacizumab 15mg/kg over 60mins (30-90mins)+cisplatin 50mg/m<sup>2</sup> IV over 60mins(30-90mins)+Paclitaxel 135mg/m<sup>2</sup> IV over 3hours
2. Topotecan 0.75mg/m<sup>2</sup> IV on day 1-3 over 30mins follow by paclitaxel 135mg/m<sup>2</sup> over 3hours+Bevacizumab 15mg/kg IV over 60mins
3. Cisplatin 50mg/m<sup>2</sup> IV+Paclitaxel 135 mg/m<sup>2</sup> IV over 3hours

## **CHEMOTHERAPY REGIMENS FOR RECURRENT OR METASTATIC CERVICAL CANCER**

### **(References)**

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- <sup>2</sup>Monk BJ, Sill MW, McMeekin DS, et al. Phase III trial of four cisplatin-containing doublet combinations in stage IVB, recurrent, or persistent cervical carcinoma: A Gynecologic Oncology Group Study. J Clin Oncol 2009;27:4649-4655.
- <sup>3</sup>Moore DH, Blessing JA, McQuellon RP, et al. Phase III study of cisplatin with or without paclitaxel in stage IVB, recurrent, or persistent squamous cell carcinoma of the cervix: a gynecologic oncology group study. J Clin Oncol 2004;22:3113-3119.
- <sup>4</sup>Moore KN, Herzog TJ, Lewin S, et al. A comparison of cisplatin/paclitaxel and carboplatin/paclitaxel in stage IVB, recurrent or persistent cervical cancer. Gynecol Oncol 2007;105:299-303.
- <sup>5</sup>Kitagawa R, Katsumata N, Shibata T, et al. A randomized, phase III trial of paclitaxel plus carboplatin (TC) versus paclitaxel plus cisplatin (TP) in stage IVb, persistent or recurrent cervical cancer: Japan Clinical Oncology Group study (JCOG0505) [abstract]. J Clin Oncol 2012;30(Suppl 15):Abstract 5006.
- <sup>6</sup>Long HJ, 3rd, Bundy BN, Grendys EC, Jr., et al. Randomized phase III trial of cisplatin with or without topotecan in carcinoma of the uterine cervix: a Gynecologic Oncology Group Study. J Clin Oncol 2005;23:4626-4633.
- <sup>7</sup>Brewer CA, Blessing JA, Nagourney RA, et al. Cisplatin plus gemcitabine in previously treated squamous cell carcinoma of the cervix. Gynecol Oncol 2006;100:385-388.
- <sup>8</sup>Weiss GR, Green S, Hannigan EV, et al. A phase II trial of carboplatin for recurrent or metastatic squamous carcinoma of the uterine cervix: a Southwest Oncology Group study. Gynecol Oncol 1990;39:332-336.
- <sup>9</sup>Kudelka AP, Winn R, Edwards CL, et al. An update of a phase II study of paclitaxel in advanced or recurrent squamous cell cancer of the cervix. Anticancer Drugs 1997;8:657-661.

**Ca of cervix : High risk factor:**

**1.Deep myometrium invasion**

**2.Tumtor size  $\geq$ 4cm**

**3.Non-squamous history**

**4.Parametrium involvement**

**5.Pelvic lymph node metastasis**

## 二、放射治療政策

(一) 治癒性放射治療(definitive curative radiotherapy alone)：包括全骨盆腔體外放射治療 (whole pelvis external beam radiation therapy) 及局部劑量追加[腔內近接治療(intracavitary radiotherapy, ICRT; intracavitary brachytherapy, ICBT) , 或強度調控放射治療(intensity modulated radiation therapy)。

適應症 (indications)：

1. 早期之子宮頸癌 (IA2, IB1,或 IIA 且腫瘤直徑小於四公分)，不宜或不願手術治療者
2. IB2,或 IIA 且腫瘤直徑大於四公分者(可考慮合併以 cisplatin 為主之化學治療)
3. IIB 以上較晚期之子宮頸癌(可考慮合併以 cisplatin 為主之化學治療)

(二) 術後放射治療(postoperative radiotherapy)：包括全骨盆腔體外放射治療及局部劑量追加[陰道內近接放射治療 (intravaginal radiotherapy, IVRT or intravaginal brachytherapy, IVBT), 或強度調控放射治療(intensity modulated radiation therapy)。

適應症 (indications)：

早期子宮頸癌 (IA,IB1,或 IIA 腫瘤直徑小四公分者)，手術治療後，病理報告有下列情況者，建議考慮放射治療。

1. 深層基質受侵犯(deep stromal invasion)
2. 淋巴血管受侵犯(lymphovascular invasion)
3. 子宮頸旁組織受侵犯(parametrial invasion)
4. 手術切除邊緣發現癌細胞(positive surgical margin)
5. 骨盆腔淋巴腺轉移(positive pelvic nodes)

(三) 緩解性放射治療：針對第 IV 期病患之轉移部位(如骨骼、腦等部位)施行緩解性放射治

### 三、局部劑量追加：依病患病情與意願選擇下列技術

#### (一) 腔內近接放射治療 (intracavitary radiotherapy, ICRT or intracavitary brachytherapy, ICBT)

採高劑量率後荷式近接治療(high-dose-rate afterloading brachytherapy)

#### (二) 單純近接治療 (brachytherapy alone) 可用來治療分期為 IA1 或一些分期為 IA2 的子宮頸癌

#### (三) 較晚期之子宮頸癌，如 Bulky IB,IIB,IIIA,IIIB 則必需先給予 45-50 Gy/25 fractions/5 weeks 之體外全骨盆放射治療

#### (四) Point A 劑量

1. 單一近接治療為每分次(fraction) 7 格雷(Gy)，一週二分次，總共七分次。
2. Bulky IB,IIB,IIIA,IIIB 之體外全骨盆放射治療後近接治療為每分次(fraction) 5-6 格雷(Gy)，一週二分次，總共 5-6 分次

#### 四、陰道內近接放射治療 (**intravaginal radiotherapy, IVRT or intravaginal brachytherapy, IVBT**)

- (一) 適用於手術陰道切除(vaginal cuff)安全邊緣不足或術後放射治療後陰道仍有殘餘腫瘤之病人
- (二) 劑量為陰道黏膜下五毫米(5mm)每分次(fraction) 4-5 格雷(Gy)，一週二分次，總共四分次。

#### 五、強度調控放射治療 (**intensity modulated radiation therapy**)

完成全骨盆放射治療者，計畫標靶體積(planning target volume, PTV)應包括 GTV 外加 0.5- 1.0 公分邊界進行局部劑量追加，再給予劑量應為 25-27Gy / 14-15 分次。

六、Concurrent Chemoradiotherapy : External irradiation 49Gr with 29 fraction +Cisplatin 40mg/m<sup>2</sup>/wk for 6 weeks.

參考資料:

- 1.Monk BJ, Sill MW, McMeekin DS, et al. Phase III trial of four cisplatin-containing doublet combinations in stage IVB, recurrent, or persistent cervical carcinoma:A Gynecologic Oncology Group Study. J ClinOncol 2009;27:4649-4655.
- 2.Moore DH, Blessing JA, McQuellon RP, et al. Phase III study of cisplatin with or without paclitaxel in stage IVB, recurrent, or persistent squamous cell carcinoma of the cervix: a gynecologic oncology group study. J ClinOncol. 2004;22:3113-3119.
- 3.Moore KN, HerzogTJ, Lewin S, et al.Acomparison of cisplatin/paclitaxel and carboplatin/paclitaxel in stage IVB, recurrent or persistent cervical cancer.GynecolOncol2007;105:299-303.
- 4.Long HJ, 3rd, Bundy BN, Grendys EC, Jr., et al. Randomized phase III trial of cisplatin with or without topotecan in carcinoma of the uterine cervix: a Gynecologic Oncology Group Study. J ClinOncol 2005;23:4626-4633.
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- 6.Weiss GR, Green S, Hannigan EV, et al.Aphase II trial of carboplatin for recurrent or metastatic squamous carcinoma of the uterine cervix: a Southwest Oncology Group study. GynecolOncol 1990;39:332-336.
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- 8.NCCN clinical practice guideline in oncology-Cervical cancer (V.1.2012)
- 9.國家衛生研究院子宮頸癌篩檢及治療共識(2000 年 10 月再版)
- 10.高雄榮民總醫院子宮頸癌放射治療指引(2009 年)
- 11.佛教慈濟綜合醫院子宮頸癌放射治療標準政策與執行政序(2009 年)
- 12.International Commission on Radiation Units and Measurements. ICRU Report No 50:Prescribing, Recording and Reporting Photon Beam Therapy. Bethesda, MD: ICRU Publications 1993.
- 13.International Commission on Radiation Units and Measurements. ICRU Report No 62: Prescribing, Recording and Reporting Photon Beam Therapy (Supplement to ICRU Report 50).Bethesda, MD: ICRU Publications 1999.

	FIGO	PRIMARY TUMOR (T)
<b>TX</b>		<b>Primary tumor cannot be assessed</b>
<b>T0</b>		<b>No evidence of primary tumor</b>
<b>Tis</b>	<b>*</b>	<b>Carcinoma in situ (preinvasive carcinoma)</b>
<b>T1</b>	<b>I</b>	<b>Cervical carcinoma confined to uterus (extension to corpus should be disregarded)</b>
<b>T1a**</b>	<b>IA</b>	<b>Invasive carcinoma diagnosed only by microscopy. Stromal invasion with a maximum depth of 5.0 mm measured from the base of the epithelium and a horizontal spread of 7.0 mm or less. Vascular space involvement, venous or lymphatic, does not affect classification</b>
<b>T1a1</b>	<b>IA1</b>	<b>Measured stromal invasion 3.0 mm or less in depth and 7.0 mm or less in horizontal spread</b>
<b>T1a2</b>	<b>IA2</b>	<b>Measured stromal invasion more than 3.0 mm and not more than 5.0 mm with a horizontal spread 7.0 mm or less</b>
<b>T1b</b>	<b>IB</b>	<b>Clinically visible lesion confined to the cervix or microscopic lesion greater than T1a/IA2</b>
<b>T1b1</b>	<b>IB1</b>	<b>Clinically visible lesion 4.0 cm or less in greatest dimension</b>
<b>T1b2</b>	<b>IB2</b>	<b>Clinically visible lesion more than 4.0 cm in greatest dimension</b>
<b>T2</b>	<b>II</b>	<b>Cervical carcinoma invades beyond uterus but not to pelvic wall or to lower third of vagina</b>
<b>T2a</b>	<b>IIA</b>	<b>Tumor without parametrial invasion</b>
<b>T2a1</b>	<b>IIA1</b>	<b>Clinically visible lesion 4.0 cm or less in greatest dimension</b>
<b>T2a2</b>	<b>IIA2</b>	<b>Clinically visible lesion more than 4.0 cm in greatest dimension</b>
<b>T2b</b>	<b>IIB</b>	<b>Tumor with parametrial invasion</b>
<b>T3</b>	<b>III</b>	<b>Tumor extends to pelvic wall and/or involves lower third of vagina, and/or causes hydronephrosis or non-functioning kidney</b>
<b>T3a</b>	<b>IIIA</b>	<b>Tumor involves lower third of vagina, no extension to pelvic wall</b>
<b>T3b</b>	<b>IIIB</b>	<b>Tumor extends to pelvic wall and/or causes hydronephrosis or non-functioning kidney</b>
<b>T4</b>	<b>IVA</b>	<b>Tumor invades mucosa of bladder or rectum, and/or extends beyond true pelvis (bullous edema is not sufficient to classify a tumor as T4)</b>
		<b>*FIGO staging no longer includes Stage 0 (Tis)</b>
		<b>** All macroscopically visible lesions—even with superficial invasion—are T1b/IB</b>

	FIGO	REGIONAL LYMPH NODES (N)
NX		Regional lymph nodes cannot be assessed
N0		No regional lymph node metastasis
N1	IIIB	Regional lymph node metastasis
DISTANT METASTASIS (M)		
	FIGO	
M0		No distant metastasis (no pathologic M0; use clinical M to complete stage group)
M1	IVB	Distant metastasis (including peritoneal spread, involvement of supraclavicular or mediastinal lymph nodes, lung, liver, or bone)

STAGE			
GROUP	T	N	M
Stage 0*	Tis	N0	M0
Stage I	T1	N0	M0
Stage IA	T1a	N0	M0
Stage IA1	T1a1	N0	M0
Stage IA2	T1a2	N0	M0
Stage IB	T1b	N0	M0
Stage IB1	T1b1	N0	M0
Stage IB2	T1b2	N0	M0
Stage II	T2	N0	M0
Stage IIA	T2a	N0	M0
Stage IIA1	T2a1	N0	M0
Stage IIA2	T2a2	N0	M0
Stage IIB	T2b	N0	M0
Stage III	T3	N0	M0
Stage IIIA	T3a	N0	M0
Stage IIIB	T3b T1-3	Any N N1	M0 M0
Stage IVA	T4	Any N	M0
Stage IVB	Any T	Any N	M1
*FIGO no longer includes Stage 0 (Tis)			
Stage unknown			