The Mediastinum

SAN NATHO

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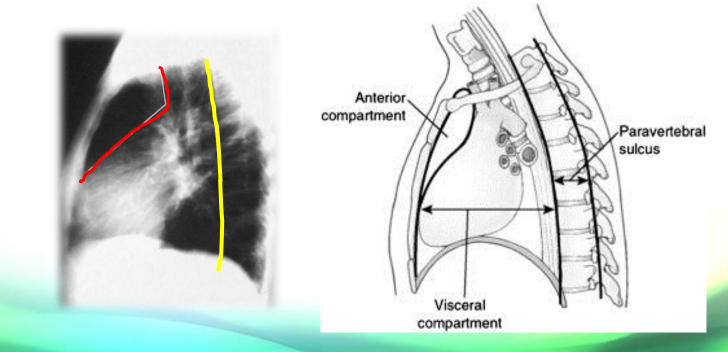


Contents

- Anatomy
- Non invasive and invasive Investigations
- Mediastinal infection
- Primary mediastinal tumors and syndromes associated with mediastinal lesions
- Mediastinal cysts

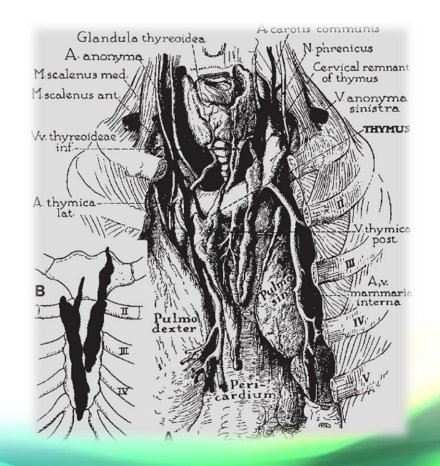


- Subdivision
 - Ant compartment
 - > Middle or visceral compartment
 - Post compartment or paravertebral sulcus





- Ant compartment
 - > Thymus
 - Internal mammary vessels
 - > Lymph nodes
 - Connective tissue with fat



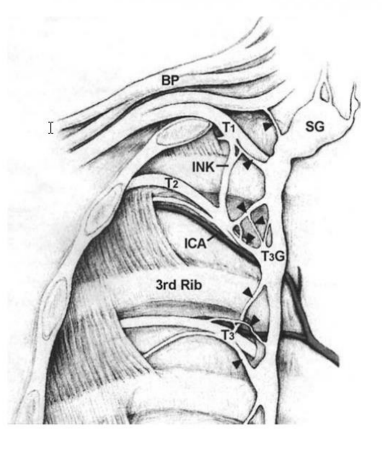
PAU 부산대학교병원

- Middle compartment
 - Pericardium, heart, and great vessels
 - Trachea, proximal portions of st esophagus
 - > Extensive lymphatic tissues
 - Vagus and phrenic nerves and and fibers
 - Supra-aortic and para-aortic bo
 - Thoracic duct, the proximal por
 - Connective tissue and fat





- Post compartment
 - > Proximal portions of the interco
 - Thoracic spinal ganglions, syr
 branches
 - Connective and lymphatic tiss

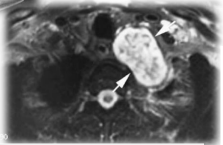




Non invasive Investigations

- Chest plain film including lateral view
- Chest CT
- Chest MRI
- Radionuclide studies
- Mediastinal tumor markers

TABLE 168-2 Serum or Urine Markers of Mediastinal Tumors		
Mediastinal location	Tumor	Serum marker
Anterior compartment	Yolk sac tumor ^a	AFP, LDH
•	Embryonal carcinoma	LDH, TRA-1–60, CD30, β -HCG ^b
	Choriocarcinoma	β -HCG, LDH
	Seminoma	PLAP, LDH, NSE, β -HCG ^b
	Thymoma	None
	Thymic carcinoma	None
	Thymic carcinoid	ACTH, chromogranin, NSE
	Thymic small cell carcinoma	Bombesin, NCAM, NSE
	Parathyroid adenoma	PTH, chromogranin
Visceral and posterior compartments	Pheochromocytoma, neuroblastoma, and ganglioneuroblastoma	Urine ^c and plasma ^d catecholamines, and chromogranin, NSE





Invasive Investigations and surgical approaches

- Transcervical mediastinal LN sampling and Lymphadenectomy
 - Mediastinoscopy: extended, video-assisted
- Video-assisted thoracic surgery
- Sternotomy and Thoracotomy
- Posterior Mediastinotomy



Mediastinal infections

- Acute and chronic mediastinitis
 - > Perforation of the aero-digestive tract
 - Postoperative sternal infection and mediastinitis
 - > Descending necrotizing mediastinitis
 - Sub-acute mediastinitis
 - Fibrosing mediastinitis



Perforation of the aero-digestive tract

- Four principles of treatment
 - Eliminate source of soilage by primary repair or diversion away from the esophageal perforation
 - 2. Provide thorough and wide mediastinal drainage to control on going mediastinal suppuration occurring after primary repair or diversion. In addition, gastrostomy tube decompression should be performed to decrease gastric reflux and mediastinal soilage.
 - Appropriate antibiotics should be administered to augment host defenses, which must be effective against both gram positive and gram negative bacteria and against both aerobic and anaerobic bacteria.
 - 4. Maintain adequate nutrition.



Case

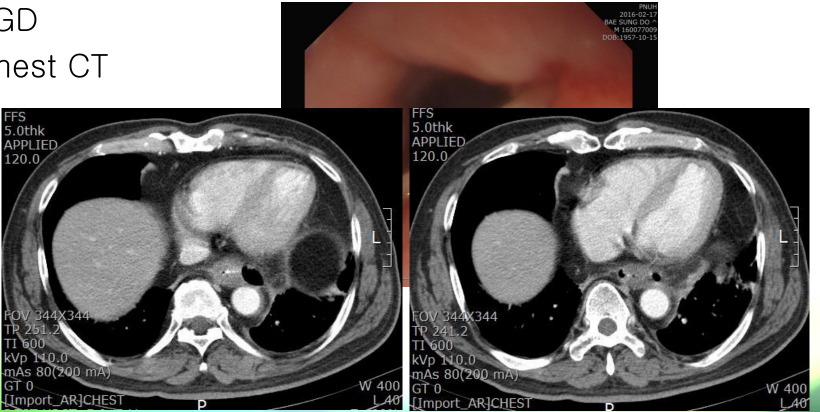
- 56/M
- 내원 수일전 매운탕 먹다가 목에 이물감 발생
- EGD •
- Chest CT •

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GT 0





Postoperative sternal infection and mediastinitis

- Risk factor
 - Sternotomy: incomplete closure
 - Tracheostomy
 - CPB duration
 - Postoperative bleeding
 - Infection
 - Low cardiac output
 - Poor general condition
 - Steroid



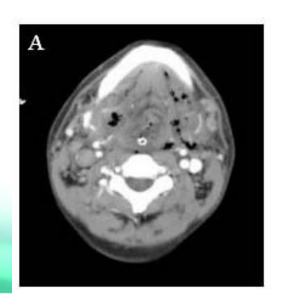
Descending necrotizing mediastinitis

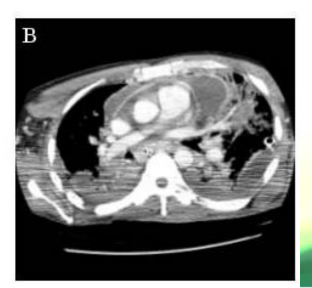
- Acute purulent mediastinitis due to oropharyngeal infection
- uncommon but still lethal form of mediastinitis
- 60 ~ 70%, secondary to odontogenic infections
- Peritonsillar abscess, Retropharyngeal and parapharyngeal abscess, Epiglottitis
- Other less common causes
 - trauma to the neck, including neck or mediastinal surgery
 - cervical lymphadenitis, endotracheal intubation



Case

- 43세 여자 환자가 고열과 전신무력감 호소
- Present illness : 최근 치통으로 충치치료를 지속적으로 받고 있 었으나 잘 조절되지 않아 발치를 하였으며 이후 고열과 전신무력감 이 심해짐
- V/S : BP 80/50, PR 120/min, BT 38.9'C
- P/Ex: 턱 아래쪽과 목 주위가 부어 있었으며 발적과 함께 열감과 동통
- Chest CT







• 진단은?

• 치료는?

• 예후는?



Sub-acute mediastinitis

- The definition of subacute mediastinitis is unclear, but this term should embrace those inflammatory processes involving the mediastinum that produce minimal to mild and evanescent symptomatology (substernal pain, fever, night sweats) and an identifiable anterior or visceral mediastinal mass by radiographic or CT examination.
- These infections most often are the result of fungal, mycobacterial, or, rarely, actinomycotic organisms.
- Such subacute infections are observed only infrequently in previously normal, healthy persons but are becoming more common in immunocompromised patients, particularly those with AIDS.



Fibrosing mediastinitis

- Fibrosing mediastinitis is an uncor resulting in the deposition and pro tissue through out the visceral cor mediastinum.
- This chronic inflammatory proce and compression of vital medias



Primary mediastinal tumors and syndromes

- Thymic tumors
- Myasthenia Gravis
- Benign LN disease
- Germ cell tumor
- Neurogenic tumors



Thymic tumors

Classification of Thymic Tumors

Epithelial cell tumors Thymoma Type A (Spindle cell, medullary) Type AB (Mixed) Type B1 (Predominate cortical, lymphocyte-rich, organoid) Type B2 (Cortical) Type B3 (Epithelial) Other types Micronodular thymoma with lymphoid stroma Metaplastic thymoma Microscopic thymoma Sclerosing thymoma Ð Combined thymoma and thymic carcinoma Thymic carcinoma Neuroendocrine cell tumors Thymic carcinoid, well differentiated Atypical thymic carcinoid, moderately differentiated Small cell carcinoma, poorly differentiated Tumors of adipose tissue Thymolipoma Thymoliposarcoma Miscellaneous tumors Thymic hemangioma Neuroblastoma and ganglioneuroblastoma Primary malignant melanoma Myoid tumor Lymphoid tumors



Thymoma

- Neoplasm of the thymus that originates in the gland's epithelial tissue.
- Incidence: 0.15 /100,000 person years (United States)
- Typically slow-growing tumors
- Spread by local extension
- Metastases are usually confined to the pleura, pericardium, or diaphragm, whereas extrathoracic metastases are uncommon.



The New World Health Organization Histologic Classification of Thymic Epithelial Tumors

Type A thymoma (medullary) Type AB thymoma (mixed) Type B thymoma^a Type B1 (organoid) Type B2 (cortical) Type B3 (epithelial) Type C (thymic carcinoma)

^aMay include combinations of B2 and B3 as well as Bl and B2.

ТА	BLE 188-7 Staging	Schemes of Thymoma	
Stage	Bergh, et al. ²¹	Masaoka, et al. ¹⁷⁹	
I	Intact capsule or growth withi the capsule	in Macroscopically, completely encapsulated; microscopically, no capsular invasion	
п	Pericapsular growth into med fat tissue	liastinal	
IIA		Macroscopic invasion into surrounding fatty tissue or mediastinal pleura	5 year survival rate
			Stage I — 94 - 100 %
IIB		Microscopic invasion into capsule	Stage II — 86 - 9 <mark>5 %</mark>
III	Invasive growth into the surro organs, intrathoracic metas or both	· · · · ·	Stage III <u>- 56 - 69 %</u>
IVA		Pleural or pericardial dissemination	Stage IV — 11 - 50 %
IVB		Hematogenous or lymphogenous	
		metastases	



Clinical presentation

- Thoracic symptoms
 - Related to the size of the tumor and its effects on adjacent organs
 - : chest pain, shortness of breath, cough, phrenic nerve palsy, superior vena cava obstruction
 - Systemic ("B") symptoms
 - : fever, weight loss, and/or night sweats

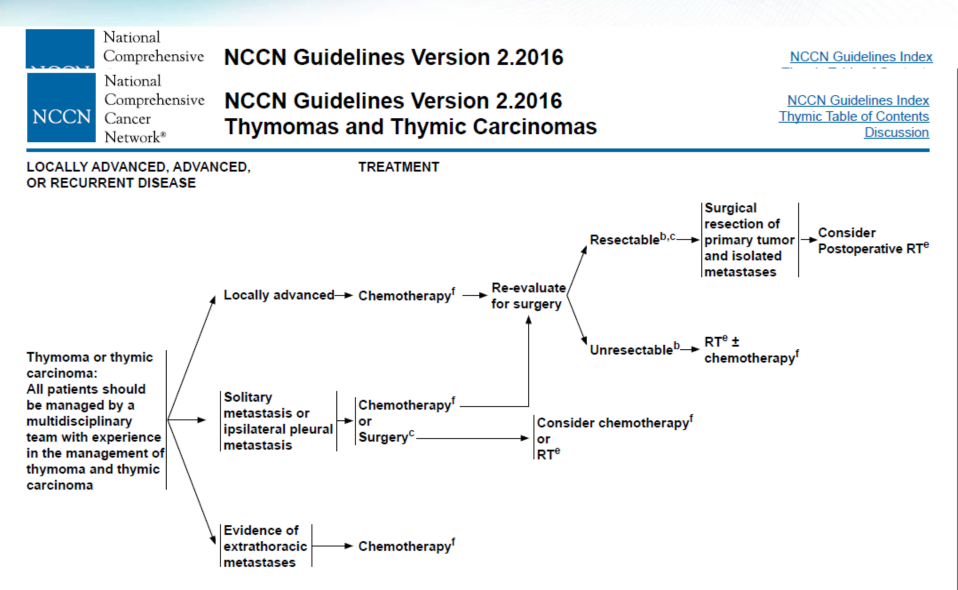


Paraneoplastic disorders

- Myasthenia gravis
- Pure red cell aplasia
- Immunodeficiency
- Thymoma-associated multiorgan autoimmunity



Treatment





Prognosis

- Thymomas
 - usually slow-growing tumors
 - presence of invasion is an important adverse prognostic marker
- The overall five-year survival : 70 %
 - 50 % with local invasion
 - 75 % without invasion
- 10-year survival : 50 %
 - 30 % with invasion
 - 60 % without invasion

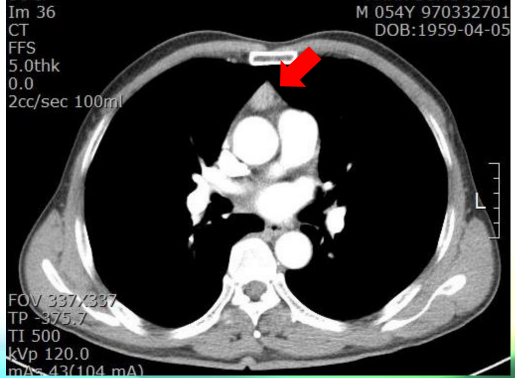


Myasthenia Gravis

- Neuromuscular junction disorder
- caused by the autoimmune destruction of the acetylcholine receptors of voluntary muscle
- Sx: diplopia, ptosis, dysphagia, weakness, fatigue
- 30 50 % of patients with thymomas have myasthenia gravis
- rare in thymic carcinoma



Case





• 진단을 위한 검사는?

• 진단은?

• 적절한 그 다음 조치는?



Benign LN disease

Benign Mediastinal Lymphadenopathies

- I. Mediastinal granulomatous disease
 - Tuberculosis
 - Fungal infection
 - Sarcoidosis
 - Silicosis
 - Wegener's granulomatosis
- II. Castleman's disease
- III. Others
 - Systemic lupus erythematosus Infectious mononucleosis Reactive lymph node hyperplasia Amyloidosis HIV–associated *Pneumocystis carinii*

Source: Machevsky MA, Kaneko M. Surgical Pathology of the Mediastinum. New York: Raven Press, 1984:174. With permission.



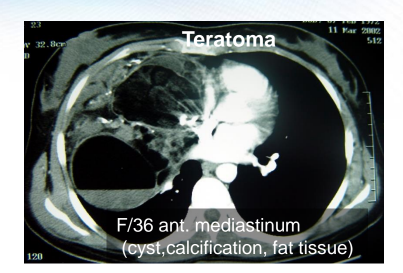
Germ cell tumor

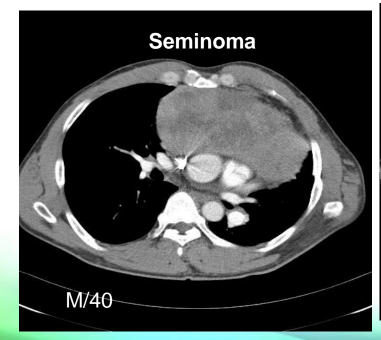
- 1. Benign germ cell tumors
- 2. Primary seminomas
- 3. Non-seminomatous malignant germ cell tumors



Mediastinal Tumor

- Anterior mediastinum
- Thymoma? Lymphoma? Teratoma? or other
- Biopsy?
- Operation?
 - When?
 - (VATS or sternotomy? thoracotomy?)
- Postop. ?





Choriocarcinoma





Classification

양성 생식세포종 (Benign GCT)	Epidermoid cyst	Teratomatous tumor
	Dermoids (dermoid cyst)	Mature teratoma
	Teratoma (mature teratoma)	Immature teratoma
		Teratoma with additional malignant component
정상피종 <mark>(Seminoma)</mark>		Non-teratomatous Tumors
비정상피종성 생식세포종 (NSGCT)	Malignant teratoma	
Choriocarcino Yolk sac carci Embryonal car		
Teratocarcino		

GCT; Germ cell tumor NSGCT; non-seminomatous germ cell tumor

Mullen & Richadson (1986) WHO (Mostofi and Sobin, 1977) - Mediastinal Germ Cell tumors(1977a) Testicular GCT (British testicular Tumor Panel; 1953, 1973, 1976)



Incidence

5-10%

(extra-gonadal, mediastinum)

15% (85% benign)

25% (children, 대부분 benign)

42 (10%) (50% benign)

Benign GCT (Teratoma) Shirodkar (1997)

97-98% anterior mediastinum 3-8% posterior mediastinum)

Malignant GCT

1-5% of all germ cell neoplasm3-5% of mediastinal tumors

Seminoma 50% / Non-seminomatous GCT 50%

of Germ cell tumor

of Anterior mediastinal tumors

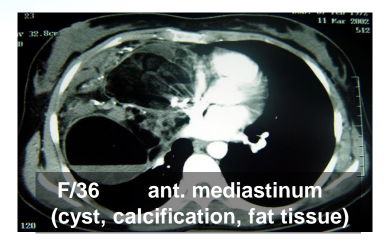
Mullen & Richardson (1986)

400 mediastinal mass

Duke Univ. medical center (1930-1982)



Benign Germ Cell Tumors





F/21 ant. mediastinum (cystic)



M/45 ant. mediastinum (cyst with fat tissue)



F/4 ant. Mediastinum (calcification, cyst with fat)



Three primodial layers

- Ectoderm; skin, hair
- Mesoderm; bone, fat, muscle
- Endoderm; respiratory epithelium, GIT
- Mature cells or tissues Mature teratoma
- Less well-differentiated tissues **Immature**
 - Infant; behave similarly to mature teratoma
 - Older patient; more aggressive (malignant teratoma)



Ruptured mediastinal Teratoma

• Incidence of spontaneous rupture

up to 36% into lung & bronchial tree,

pleural space, pericardial space, great vessels

- Hypothesis of rupture
 - Autolysis : most compelling cause, digestive enzymes (pancreatic tissue, salivary gland tissue)
 - Chemical inflammation : sebaceous gland secretions
 - Ischemia : rapid enlargement
 - Pressure necrosis : thinning of the cyst wall d/t secretions
 - Infection : tumor wall fragile(pulmonary or hematologic)

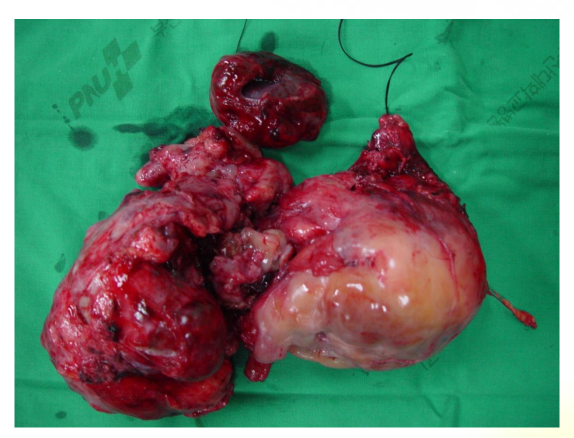
Spontaneous rupture of benign mature teratomas of the mediastinum Am J Roentgenol. 1998 Feb:170(2):323-328 Teratoma with malignant transformation in the ant. Mediastinum Korean J Radiol. 2000 Jul-Sep:1(3):162-4



10 months old male Huge mediastinal mass with pleural effusion



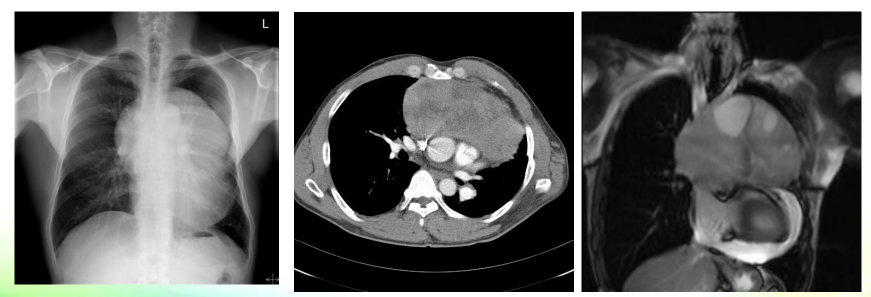






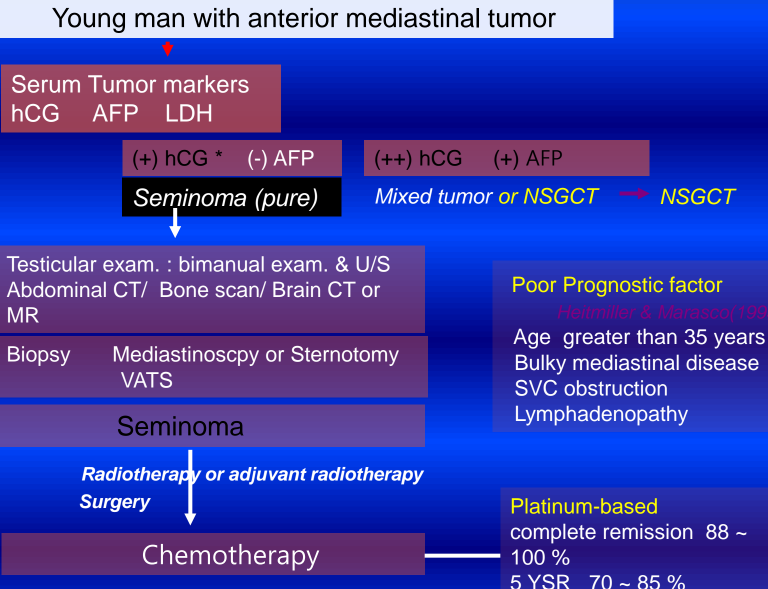
Seminoma

- Second common mediastinal GCT / TMC malignant mediastinal GCT
- 3 rd ~ 5 th decade men, white men predominant
- Slow-growing tumors with lobular appearance including necrosis, hemorrhage
 - encapsulation half of time, calcification infrequently



40/M Seminoma

Seminoma





International Germ Cell Cancer Collaboration Group J Clin Oncol 1977

Good Prognosis		
Any Primary site	90% of seminomas	
No NPVM	5 year PFS 82 %	
Normal AFP, hCG, LDH	5 YSR 86 %	
Intermediate Prognosis		
Any Primary site	10% of seminomas	
NPVM (liver, bone, brain)	5 year PFS 67 %	
Normal AFP, hCG, LDH	5 YSR 72 %	

PFS progression free survival

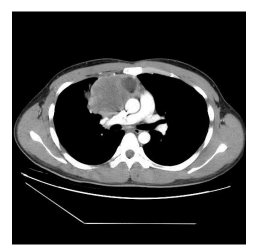


Non-seminomatous GCT

- Potentially curable with surgery
- Exclusively in young adult, men (fewer than 30 cases reported in women)
- Rapid local growing tumors with early metastasis (85-90% at diagnosis)
- In-homogenous mass with multiple areas of necrosis & hemorrhage



30/ M Choriocarcinoma





24/ M Endodermal sinus tumor



NSGCT

Incidence	Moran & Suster (1997)	강창현 (2008)	
	229 cases	29cases	
Teratocarcinoma	41 %	9.5 %	

58% non-germ cell component (sarcoma, epithelial carcinoma)

Endodermal sinus (Yolk sac) tumor	35 %	42.9 %
Choriocarcinama	7 %	4.8 %
Embryonal carcinoma	6 %	9.5 %
Mixed	11 %	9.5 %
Unknown		23.8 %

Differ from testis origin

Pure endodermal sinus tumor, extremely rare in testis

Embryonal carcinoma, much higher in testis

Non-germ call histologies is more common in mediastinum





Tumor markers

hCG or AFP	90%
AFP with/without hCG	80%
hCG	30-35%
LDH	80-90%

APF 이 증가된 경우는 조직검사상 pure seminoma로 보인다고 해도 NSGCT와 같이 치료

hCG가 100 ng/ml 이상은 pure seminoma에서 uncommon

Differ from testis origin

Testicular NSGCT AFP & hCG equal frequency

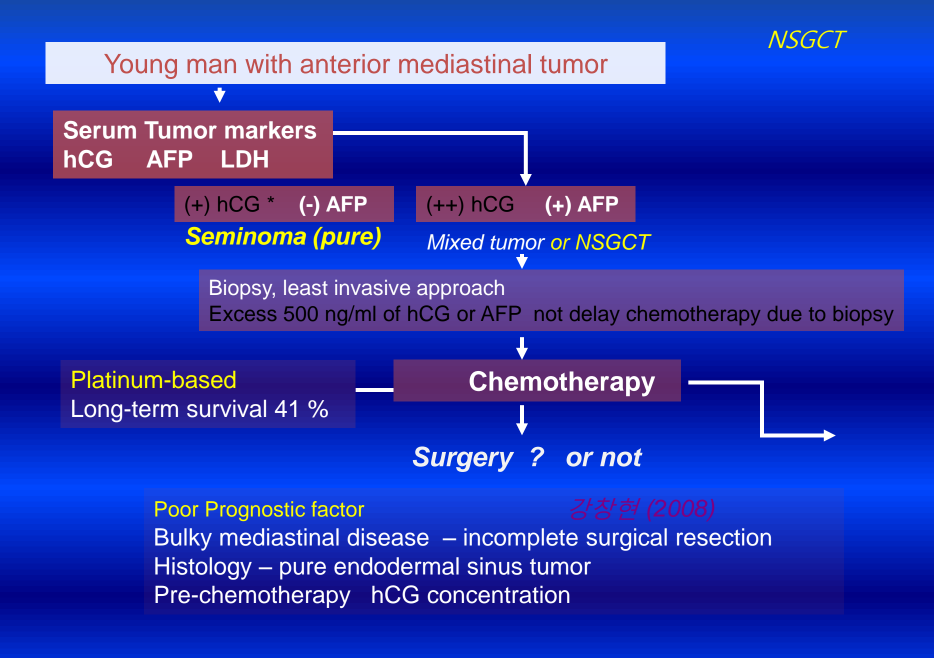


NSGCT

Associated syndromes

Hematologic malignancies

Acute non-lymphocytic leukemia Erythroleukemia Myelodysplastic syndrome		Acute lymphocytic leukemia Acute megakaryocytic leukemia Malignant histiocytosis				
				Hart	mann(2000) 2%	Median survival 5 months
				(287 me	diastinal NSGCT)	No patient more than 2 years
Idiopathic thrombocytopenia						
Hemophagocytic syndrome	single ca	ase of endodermal sinus tumor				
Klinefelter's syndrome no	ot associated with	n testicular GCT				
comn	on underlying ge	rm cell defect				



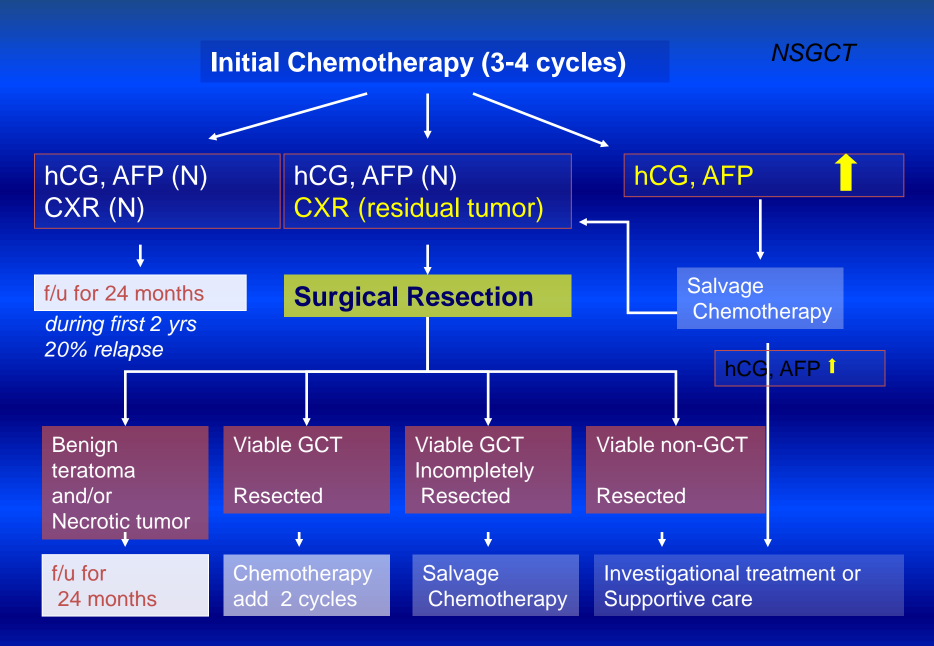


Fig 187-7 Shield, General Thoracic Surgery



NSGCT

International Germ Cell Cancer Collaboration Group J Clin Oncol 1977

Good Prognosis	AFP	hCG	LDH	non-semoi	nomas
Testis/retroperitoneal				56%	
No NPVM				5 year PFS	82 %
Good markers	< 1000	< 1000	< 1.5 x N	5 YSR	86 %
Intermediate Prognosis					
Testis/retroperitoneal				28%	
No NPVM				5 year PFS	75 %
Intermediate markers	1,000~	1,000~	1.5 x~	5 YSR	80 %
	10,000	10,000	10 x N		
Poor Prognosis					
Mediastinal primary				16%	
NPVM (liver bone, brain)				5 year PFS	41 %
Poor markers	> 10,000	> 10,000	> 10 x N	5 YSR	48 %

NPVM non-pulmonary visceral metastasis PFS progression free survival



Neurogenic tumors

TABLE 196-1	Mediastinal Neurogenic Tumors in Infants and Children			
Tumors of autonomic ganglia	Neuroblastoma	Ganglioneuroblastoma	Ganglioneuroma	
Tumors of nerve sheath origin	Schwannoma	Neurofibroma	Neurogenic sarcoma	
Tumors of neuroectodermal origin	Melanotic progonoma	Askin's tumor		
Tumors of paraganglia	Paraganglioma			



Mediastinal cyst

- Foregut cyst
- Gastroenteric and Neurenteric cyst