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Testicular cancer

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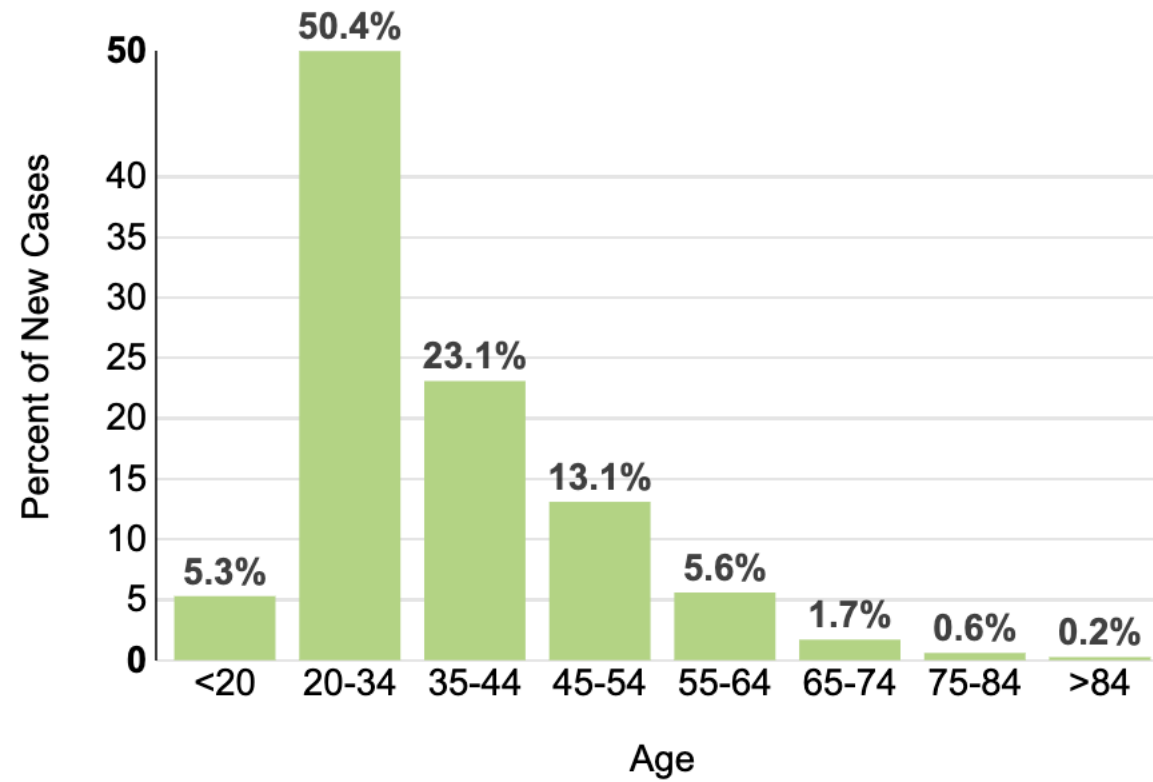
Disclosures

None

Epidemiology

- In 2021, estimated 9,470 new diagnoses
 - 440 deaths from testicular cancer
- Increasing incidence over the last several decades
 - Particularly in Hispanic Americans

Epidemiology



Epidemiology

- Risk Factors
 - Cryptorchidism (RR=10-15, absolute risk 2-3%)
 - Klinefelter's syndrome
 - Personal history (2-3% risk of contralateral 2nd primary)
 - Infertility
 - Family history (Brother RR=8-10, Father RR=4)
 - Germline CHEK2 mutations

Pathology

- Seminoma
- Non-seminoma
 - Embryonal – worse prognosis for stage I
 - Choriocarcinoma
 - Yolk sac tumor – better prognosis for Stage I
 - Teratoma
- If any histology other than seminoma → non-seminoma
- If alpha-fetoprotein is elevated → non-seminoma
- Other rare histologies – sex cord/stromal, lymphoma (>70yr)

Pathology

- Teratoma
 - Higher malignant potential in men than women or children
- Isochromosome 12p
 - Occurs in 50% of germ cell tumors
 - Excess copies of 12p can help identify some poorly differentiated carcinomas as GCT through FISH/cytogenetics

Presentation

- Painless testicular mass is pathognomonic
- Testicular pain or discomfort
- Testicular swelling
- Growth or shrinkage of testicle
- Abdominal pain/mass
- Back pain
- Gynecomastia and/or gynecodynia
- Supraclavicular and/or cervical lymphadenopathy
- Renal failure
- Lower limb edema
- Infertility

Evaluation

- H&P
- Scrotal ultrasound
- Tumor markers
 - B-HCG, AFP, LDH
- Chemistry panel
- Radical inguinal orchiectomy
- Consider sperm banking

Evaluation

- Pure seminoma
 - CT AP
 - CXR
 - CT chest if RP LAD or abnormal CXR
 - Repeat tumor markers
 - Brain MRI
 - HCG >5,000
 - Extensive lung mets
 - Symptoms

Evaluation

- Non-seminoma
 - CT CAP
 - Repeat tumor markers
 - Staging is based off post-orchietomy values
 - Brain MRI
 - HCG >5,000, AFP >10,000
 - Choriocarcinoma
 - Extensive lung mets
 - Liver mets
 - Symptoms

AFP

- Half-life 5-7 days
- Not produced by seminoma
- Can be associated with numerous cancer, but mostly hepatocellular carcinoma and non-seminomatous germ cell tumor
- AFP levels <20 ng/mL can be non-specific and treatment decisions should not be based on this alone
- Can be produced by teratoma at low levels
- May be elevated due to liver disease or hepatotoxicity (any liver regenerative state)

β HCG

- Half-life 1-3 days
- Can be made by any type of germ cell tumor
- Extremely high levels suggest choriocarcinoma
- False positives
 - Cross reactivity with luteinizing hormone
 - Can test for this by administering exogenous testosterone
 - Pituitary production in hypogonadal men
 - Marijuana consumption may lead to elevated B-HCG

Lactate Dehydrogenase

- Many conditions can elevate LDH
- Useful only for staging of disseminated disease
- The only important LDH is the level on day 1 of the first cycle of first-line chemotherapy for disseminated disease
- Treatment decisions should never be made on elevated LDH alone

Serum Tumor Markers

	Good (S1)	Intermediate (S2)	Poor (S3)
AFP (ng/mL)	<1,000	1,000-10,000	>10,000
BHCG (IU/L)	<5,000	5,000-50,000	>50,000
LDH*	<1.5x ULN	1.5-10x ULN	>10x ULN

* In practice, cutoff of >2.5-3x ULN is generally used

Staging

- Stage I
 - Limited to testis, scrotum, and spermatic cord
- Stage II
 - Metastases to retroperitoneal lymph nodes only
 - Tumor markers normal (S0) or S1
- Stage III
 - Distant metastases (including pelvic nodes)
 - RP nodal mets only but S2/S3

Risk stratification for advanced disease

	Good	Intermediate	Poor
Seminoma	Primary Site: Any Mets to nodes and/or lung	Non-pulmonary visceral mets	None
Non-seminoma	Primary Site: testis or RP Mets to nodes and/or lungs S0-1	Primary Site: testis or RP Mets to nodes and/or lungs S2	Primary site: Mediastinum Non-pulmonary visceral mets S3

Survival Based on Risk Categories for Advanced Disease

Risk Group	Percent of Patients	5 year survival
Good risk	60%	~95%
Intermediate risk	26%	~89%
Poor risk	14%	67%

International Germ Cell Cancer Collaborative Group updates. J Clin Oncol. May 2021.

Important Considerations

- Testis masses necessitate urgent workup
- Do not biopsy the testis
- Radical inguinal orchiectomy is the standard since transscrotal orchiectomy can lead to seeding of disease and increased local recurrence rates
- Discuss sperm banking prior to surgery, radiation, or chemotherapy (~20% risk of infertility)
- The testis is a sanctuary site
 - Even in patients with metastatic disease at diagnosis, radical inguinal orchiectomy should be performed, either before or after chemotherapy

Stage I Seminoma

- Active Surveillance - preferred
 - Risk of relapse~15%
 - Tumor size >4cm and rete testis involvement are risk factors for recurrence
- Adjuvant chemotherapy
 - 1-2 doses carboplatin AUC 7
 - ~2% recurrence rate
 - May decrease risk of contralateral primary
- Adjuvant radiation therapy
 - 25-30 Gy to infradiaphragmatic LNs
 - ~4% relapse rate
 - Risk of secondary cancer, GI complications, cardiovascular disease

Oliver et al, JCO 2011;29(8):957-962

Solberg et al, Ann Onc 2016;27(7):1299-1304

Aparicio et al, JCO 2011;29:4677-4681

Stage I Seminoma

- With any option, survival ~100%
- Relapse with tumor markers or measurable disease is treated as the stage at recurrence
- Caution with Stage IS
 - Generally portends occult disease
 - Consider false positive BHCG

Stage II Seminoma

- Nodes <3cm (IIA/IIB)
 - Radiation therapy or chemotherapy (BEPx3 or EPx4)
 - BEP – Bleomycin, etoposide, cisplatin; EP – etoposide, cisplatin
- Nodes >3cm (IIB/IIC)
 - Chemotherapy BEPx3 or EPx4
- No randomized trials

Stage I Non-seminoma

- Surveillance - preferred
 - 25% relapse
 - LVI, >pT2, and high % embryonal histology predictive of relapse
- Retroperitoneal lymph node dissection
 - 20% likelihood of finding residual disease
 - Unclear who benefits from adjuvant chemo – typically given for >5 nodes or >2cm in size
 - 11% risk of relapse
 - 10-20% of patient get chemotherapy
 - 10% risk of retrograde ejaculation

Stage I Non-seminoma

- Chemotherapy
 - 1-2 cycles BEP
 - 2% risk of relapse
- Stage IS – treat as advanced disease with chemotherapy

Stage II Non-seminoma

- IIA with normal markers
 - RPLND
 - 30% will be benign
 - Use of adjuvant chemo based on amount/size of nodes
 - BEP x3 or EP x4
 - If borderline LAD, consider short interval repeat imaging
- IIA with S1, IIB/IIC
 - BEP x3 or EP x4

Good Risk Disease

- BEP x3
 - Equivalent to BEP x4
 - Less cisplatin – anorexia, nausea, vomiting, neurotoxicity, ototoxicity, infertility
 - Less risk of etoposide-induced leukemia
 - Dose-dependent
- EP x4
 - EP x4 superior to EP x3
 - Bleomycin can cause pulmonary fibrosis
 - Post-chemo RPLND is more difficult after bleomycin
 - Consider in >50yr, renal insufficiency, pre-existing lung disease

De Wit et al, JCO 2001;19(6):1629-1640
Culine et al, Ann Onc 2007;18(5):917-924

BEP x3 vs EP x4

GETUG T93BP – 257 patients, 1:1 randomization

	BEP x3 (127)	EP x4 (124)	P-value
G3-4 Neutropenia	47%	62%	<0.001
G1-3 Neurotoxicity	2	7	<0.001
Adverse Events*	13%	22%	0.05
PFS	91%	86%	0.135
4yr OS	96%	92%	0.096

Indiana University Testis Cancer Database – 223 patients

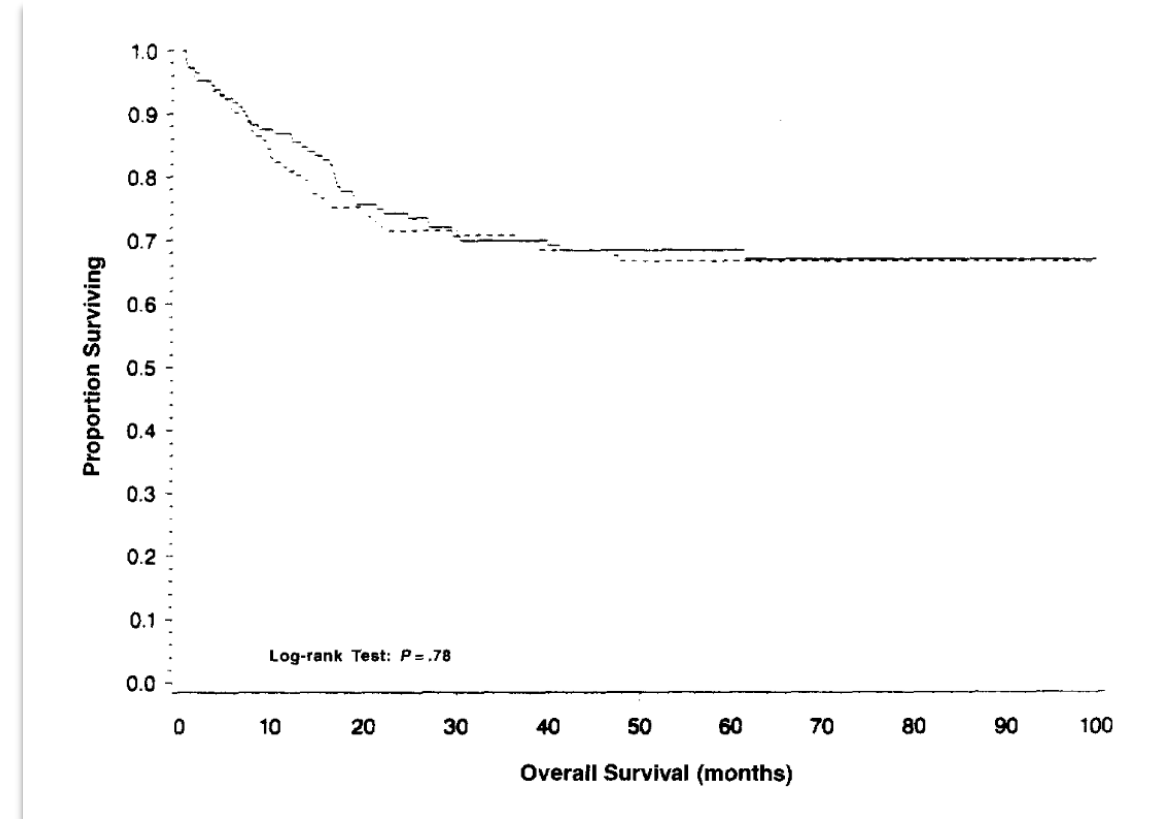
	BEP x3 (178)	EP x4 (45)	P-value
10yr OS	98%	91%	<0.01

Culine et al, Ann Onc 2007;18(5):917-924

Cary et al, Clin GU Cancer 2018;16(2)e307-e313

Intermediate and Poor Risk Disease

- BEP x4 or VIP x4
 - VP-16 (etoposide), ifosfamide, cisplatin
 - VIP for patients with concern for bleomycin pulmonary toxicity
 - Increased hematologic toxicity with VIP
 - VIP given inpatient



Nichols et al, JCO 1998;16(4)1287-1293

Post-chemotherapy management

- Seminoma
 - Most residual masses are benign
 - <3cm – 3% carcinoma (path+ or relapse)
 - >3cm – 30% carcinoma
 - Observe
 - Observe masses <3cm, resect/biopsy if >3cm
 - Observe <3cm, PET if >3cm → resect/biopsy if PET+
 - Generally wait until at least 6-8 weeks post-chemo for PET
 - Improved sensitivity and specificity

Post-chemotherapy management

- Non-seminoma
 - Resect residual masses >1cm when possible
 - Residual mass histology
 - Viable carcinoma: 10%
 - Teratoma: 40%
 - Fibrosis/necrosis: 50%
 - Teratoma needs to be removed
 - Growing teratoma syndrome
 - Malignant transformation
 - Chemo resistant

Recurrent/Relapsed Disease

- For Stage I surveillance and Stage I/II treated with RPLND or RT, treat based on stage at time of recurrence
- Post-chemo recurrence
 - Most often <2 yrs for NSGCT, <3yr for seminoma
 - Salvage chemotherapy
 - VIP x4
 - VeIP x4 (vinblastine, ifosfamide, cisplatin)
 - TIP x4 (paclitaxel, ifosfamide, cisplatin)
 - High-dose chemotherapy with autologous stem cell rescue

HDC with Autologous Stem Cell Rescue

- No benefit over standard chemotherapy for 1st line treatment
- No high-quality studies comparing HDC to standard salvage chemotherapy
- Retrospective analysis from Indiana University
 - Tandem transplant with carboplatin 700mg/m² and etoposide 750mg/m² qd x3d
 - 364 patients
 - 2yr OS 66%
 - 2nd line – 2yr PFS 63%; 3rd line – 2yr PFS 49%
- TIGER Trial – salvage chemo for HCD with TI-CE

Adra et al, JCO 2017;35(10):1096-1102

Late Relapse

- Often can be cured
- Resection is integral to the plan
- At risk for subsequent relapse

Seminoma Stage I	NSGCT Stage I	Pathological stage II*	Good-risk disease	Intermediate- or poor-risk disease
Carboplatin 1 or 2 doses	BEP x 1 or BEP x 2	BEP x 2 or EP x 2	BEP X 3 or EP X 4	BEP X 4 or VIP X 4

	Relapsed after first-line chemotherapy
Salvage Treatment	VeIP x 4 or TIP x 4 or HDCT x 2

*Pathological stage II refers to patients who had positive nodes with GCT after undergoing primary RPLND for Stage I/II disease.

Surveillance

- Clinic visit, tumor markers, imaging
 - Decrease frequency over time away from treatment
- Less intense follow-up for patient who have had systemic therapy
- Trend to using less imaging due to concern over radiation exposure
 - MRI can be used in place of CT; CXR in place of CT chest
- Consult NCCN guidelines as recommendations change frequently

Survivorship

- Cardiovascular disease risk increases ~2X
- Metabolic syndrome up to 10X risk
- Infertility
- Hypogonadism
- Erectile dysfunction – often with normal testosterone levels and may be a neuropathy
- Secondary malignancy risk increases 1.5-2X
- Contralateral primary testicular cancer – 2-3%
 - Testicular self-exam, exam at clinic visits

Survivorship

- Restrictive pulmonary disease - may be more related to cisplatin than bleomycin
- Hearing loss, tinnitus
- Peripheral neuropathy
- Renal dysfunction
- Raynaud's phenomenon

Key Points

- Affects young men and is highly curable, even with advanced disease
- Tumor markers are critical for diagnosis, staging, prognosis, treatment response, and surveillance
- For Stage I, surveillance is preferred
- Chemo-sensitive: don't dose reduce or delay!
- High-dose chemotherapy with autologous stem cell rescue can be curative
- Patients can have significant long-term side effects from treatment

Things to Remember for the Boards

- Diagnosis
 - Seminoma vs non-seminoma
 - Staging
 - Risk stratification for Stage III disease
- Use of serum tumor markers for staging, prognosis, treatment response, and surveillance
 - Know the half-lives of AFP (5-7 days) and BHCG (1-3 days) and causes of false positives
- Treatment options by stage and risk

Things to Remember for the Boards

- Management of residual masses
 - PET for seminoma >3cm
 - Resection for NSGCT
 - Risk of teratoma and persistent disease
- Complications and toxicity of treatment