

Optometric Education Consultants

**Thyroid and Thyroid Eye Disease
Clinical Pearls and Innovations for 2023**

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Phoenix, AZ
Saturday, April 15, 2023

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Disclosures- Greg Caldwell, OD, FFAO
All relevant relationships have been mitigated

- Lectured for: Alcon, Allergan, Aerie, B&L, BioTissue, Kala, Maculogix, Optovue, RVL, Heru, Santen
- Disclosure: Receive speaker honorariums
- Advisory Board: Allergan, Alcon, Dompe, Eyenovia Tarsus, Visus
- I have no direct financial or proprietary interest in any companies, products or services mentioned in this presentation
- Disclosure: Non-salaried financial affiliation with Pharmanex
- Envolve: PA Medical Director, Credential Committee
- Healthcare Registries – Chairman of Advisory Council for Diabetes and AMD
- The content of this activity was prepared independently by me - Dr. Caldwell
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Financial Obligations

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My Practice

I am a clinician first then a scientist

- Some are scientists first then clinician
- I need to simplify for patient and patient care.
- Science is great, but not good if there isn't a clinical application.
- Some lectures are science based without clinical application.
- My lecture will be a hybrid. Showing clinical applications of the science

It is wonderful to have someone who's juggling so many aspects of optometry (scientific, clinical experience, teacher & lecturer). It is refreshing and very informative. -Sarah

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Key Tenants of Aging, Performance and Vitality

- Oxidative Stress / Inflammation
- Hormonal Balance
- Stress Hormones
- Glucose / Insulin Regulation
- GUT integrity and microbiome diversity
- Immune Balance
- Environmental Exposure/Burden
- Individuality

Credit to: James LaVaila, RPh, CCN

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A4M FEST 2022 25th ANNUAL WORLD CONGRESS
FRIDAY DECEMBER 9th
10:30AM - 6:00PM

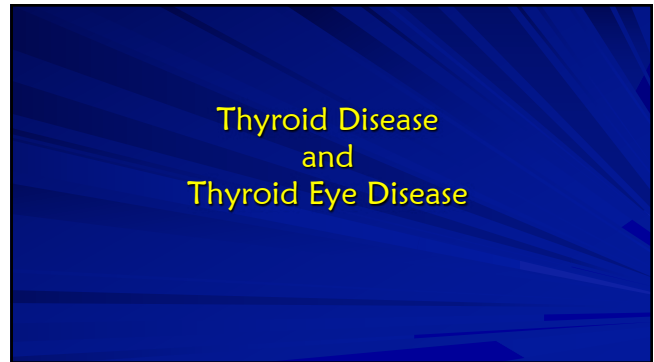
Mitochondria
The Importance of Power Density

MEM - EYE
Mitochondrial health is key to healthy aging

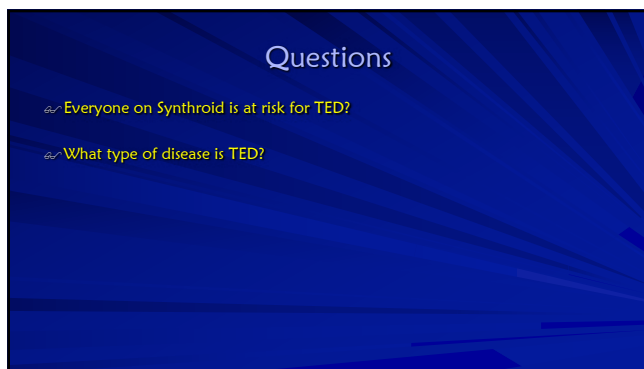
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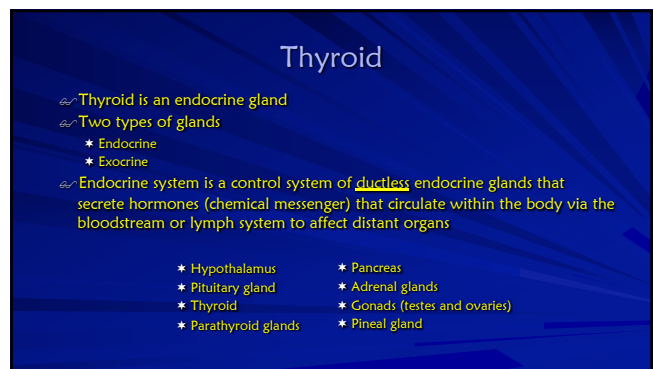
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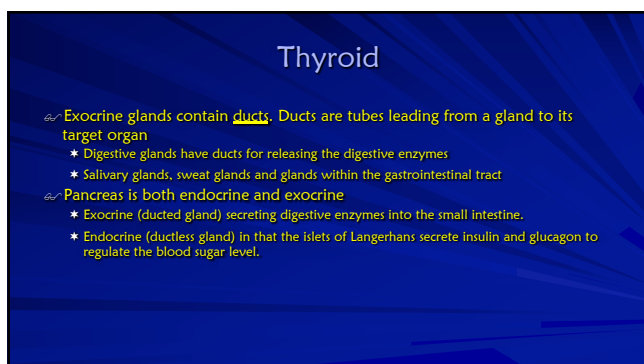
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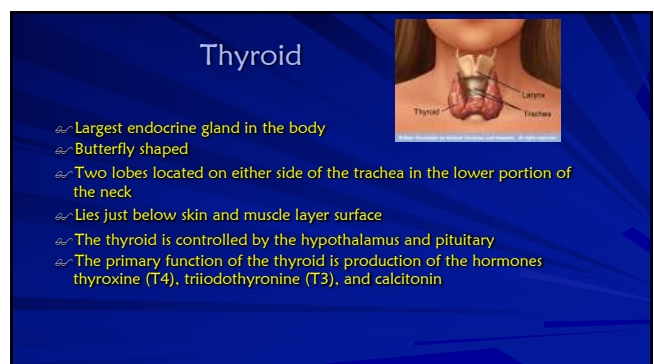
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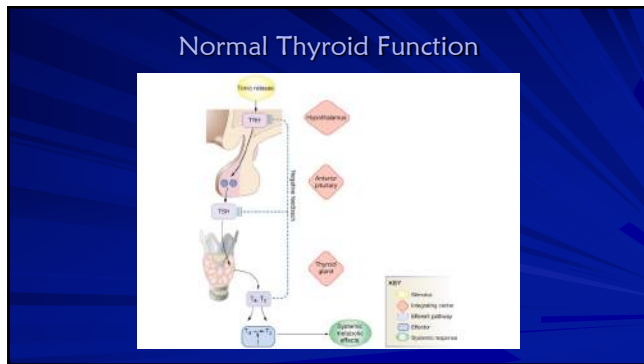
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Thyroid Dysfunction

~ What is the most common cause of thyroid dysfunction?

- Cancer
- Surgically induced
- Medication toxicity or side effect
- Pregnancy
- Autoimmune disease

~ In autoimmune disease the body typically produces _____ that attacks itself, this can be systemic or organ specific

- Antibodies, immunoglobulins

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Thyroid Dysfunction

- ~ Primary = Thyroid gland
- ~ Secondary = Pituitary failure
- ~ Tertiary = Hypothalamic

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Antibodies of Thyroid Dysfunction

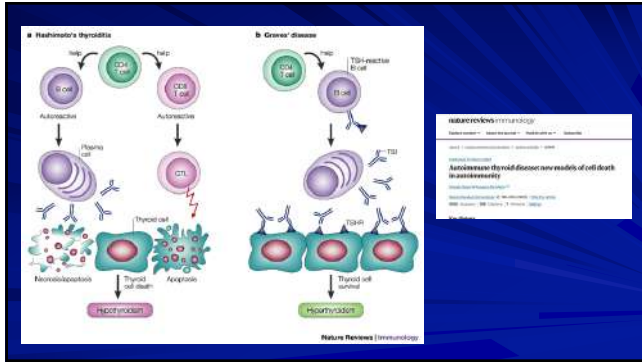
- ~ TSH Receptor Antibodies
 - * Stimulating TSH receptor antibody
 - Thyroid Stimulating Immunoglobulin (TSI)
 - * Thyroid blocking antibody (TBAb)
- ~ Thyroid Peroxidase Antibodies (TPOAb)
 - * TPO is found in thyroid follicle cells where it converts the thyroid hormone T4 to T3
 - * TPOAb contributes to thyroid cellular destruction

~ Most autoimmune thyroid dysfunctions have a combination of thyroid antibodies, however depending on which AB is more abundant results in the outcome of the disease

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Ninja Nerd Science
YouTube

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Complement factor H in AMD: Bridging genetic associations and pathobiology

Abstract: Age-Related Macular Degeneration (AMD) is a complex multifactorial disease characterized in its early stages by drusen-like accumulations in Bruch's Membrane (BM), seen on histologic examination, and in its late forms by macrophage-mediated (Pam7) or smooth muscle atrophy of the Bruch's Membrane (BM) (Pam7) and hyper-Pam7. Genetic studies have strongly supported a link between the alternative complement pathway, in particular the membrane protein factor H (CFH), and development of AMD. However, the functional significance of the CFH 1480T polymorphism remains elusive. In this review, we discuss the role of CFH in AMD pathogenesis and its potential as a therapeutic target.

Complement Cascade Effects in AMD

CFH	C3a	C5a
<ul style="list-style-type: none"> Associated with drusen formation Associated with drusen formation Associated with drusen formation 	<ul style="list-style-type: none"> Associated with drusen formation Associated with drusen formation Associated with drusen formation 	<ul style="list-style-type: none"> Associated with drusen formation Associated with drusen formation Associated with drusen formation

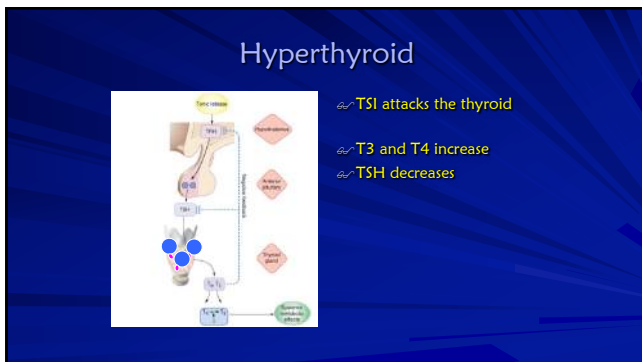
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April 27, 2021 – January 26, 2022 (9 months)

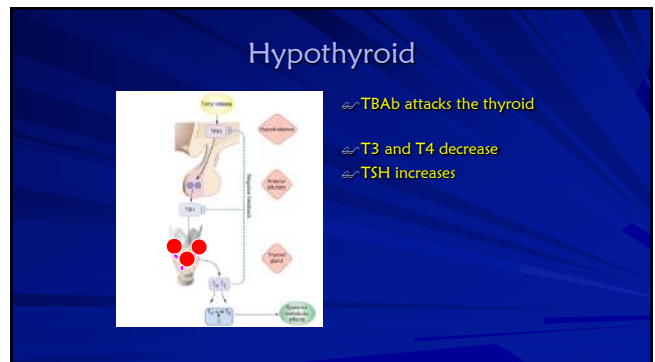
Razer Comparison Report

Melanie Clemmons, OD
May 20, 2022 AACO Nashville

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Thyroid Dysfunction

<p>Hyperthyroidism (Thyrotoxicosis)</p> <ul style="list-style-type: none"> ~ Primary-autoimmune <ul style="list-style-type: none"> * Graves <ul style="list-style-type: none"> □ Graves-Basedow or von Basedow's ~ Secondary/Tertiary <ul style="list-style-type: none"> * Excess thyroid medication for treatment of hypo or goiter * Toxic multinodular goiter * Toxic adenoma * Excess iodine * Thyroiditis (inflammatory induced) * Excess hormone production ectopic tissue * Thyroid carcinoma 	<p>Hypothyroidism (most common organ-specific autoimmune disorder)</p> <ul style="list-style-type: none"> ~ Primary-autoimmune <ul style="list-style-type: none"> * Chronic autoimmune thyroiditis <ul style="list-style-type: none"> □ Hashimoto's thyroiditis * Autoimmune atrophic thyroiditis <ul style="list-style-type: none"> □ Primary myxedema □ Opposite of Graves disease * Postpartum thyroiditis ~ Secondary/Tertiary <ul style="list-style-type: none"> * Lithium medication * Pregnancy * Surgically induced * Disorders of the pituitary gland or hypothalamus
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GRAVE'S (Hyperthyroidism)

- ~ A multisystem disorder consisting of a triad
 - * Hyperthyroidism with diffuse hyperplasia of the thyroid gland
 - * Infiltrative demopathy
 - * Infiltrative ophthalmopathy
- ~ Prevalence:
 - * 20-40 year old female (F:M = 7:1)
 - * Genetic link
- ~ Etiology:
 - * Autoimmune disease: hypersensitivity reaction with thyroid stimulation by the circulation of abnormal thyroid-stimulating immunoglobulins (TSI)

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Hashimoto's Thyroiditis (Hypothyroidism)

- ~ The most common cause of hypothyroidism in the United States
- ~ It is named after the first doctor who described this condition, Dr. Hakaru Hashimoto, in 1912
- ~ Autoimmune disease
- ~ Goiter formation
- ~ 5-10 times more common in women than in men
- ~ The underlying cause of the autoimmune process still is unknown
 - * Anti-TPO ab and Anti-TB recp ab present

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Autoimmune atrophic thyroiditis (Hypothyroidism)

- ~ Atrophic thyroiditis is similar to Hashimoto's thyroiditis
- ~ A goiter is not present

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Postpartum Thyroiditis (Hypothyroidism)

- ~ These women develop antibodies to their own thyroid during pregnancy, causing an inflammation of the thyroid after delivery

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Systemic Manifestations of Hyperthyroid (Primary or Secondary)

<p>~ Symptoms</p> <ul style="list-style-type: none"> * Nervousness * Heat intolerance * Sweating * Fatigue * Palpitation * Insomnia * Early waking * Alopecia * Vitiligo * Brittle nails 	<p>~ Signs</p> <ul style="list-style-type: none"> * Sweating * Muscle Weakness * Emotionally labile * Tremor * Tachycardia * Arrhythmia * Hypertension * Britk tendon reflex * Diabetes * ↑Triglycerides & Ca, ↓CHO * Microcytic anemia * Possible goiter * Myxedema
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Systemic Manifestations of Hypothyroid (Primary or Secondary)

- ~ Symptoms
 - * Cold intolerance
 - * Weakness
 - * Reduced energy
 - * Lethargy
 - * Muscle cramps
 - * Constipation
 - * Increased sleeping
 - * Weight gain
 - * Reduced appetite
 - * Joint stiffness
- ~ Signs
 - * Cool, scaling skin
 - * Puffy hands and face
 - * Deep voice
 - * Myotonia
 - * Delirium
 - * Bradycardia
 - * Slow reflexes
 - * Obesity
 - * Hypothermia
 - * Myxedema

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Thyroid Eye Disease (TED)

- ~ Other names used
 - * Grave's disease
 - * Grave's ophthalmopathy
 - * Grave's orbitopathy
 - * Exophthalmos in Graves Disease
 - * Thyroid Associated Orbitopathy (TAO)
 - * Thyroid Orbitopathy
 - * Ophthalmic Graves Disease
 - * Inflammatory Eye Disease
 - * Endocrine Orbitopathy

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Why is this so confusing?

- ~ Thyroid Eye Disease
 - * Is often seen in conjunction with Graves' Disease (hyperthyroid)
 - * Is seen in people with no other evidence of thyroid dysfunction
 - * Is seen in patients who have Hashimoto's Disease (hypothyroid)
- ~ Most thyroid patients, however, will not develop thyroid eye disease

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Why is this so confusing?

- ~ The eye symptoms usually occur at the same time as the thyroid disease
 - * However they may precede or follow the obvious symptoms of the thyroid abnormality
- ~ The incidence of thyroid eye disease associated with thyroid dysfunction is higher and more severe in smokers
 - * There is no way to predict which thyroid patients will be affected

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Why is this so confusing?

- ~ While eye disease may be brought on by thyroid dysfunction
 - * Successful treatment of the thyroid gland does not guarantee that the eye disease will improve
 - * No particular thyroid treatment can guarantee that the eyes will not continue to deteriorate
 - * Once inflamed, the eye disease may remain active from several months to as long as three years
 - * There may be a gradual or, in some cases, a complete improvement

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Thyroid Eye Disease

- ~ Commonly known as Graves' ophthalmopathy
- ~ About 80% of all patients with TED have the autoimmune hyperthyroid disorder known as Graves' disease
- ~ Another 10% of all cases are seen in patients with autoimmune hypothyroidism, either Hashimoto's thyroiditis, atrophic thyroiditis or Hashitoxicosis
- ~ Another 10% of all cases are seen in people with normal thyroid function
 - * When thyroid function is normal, the eye condition is referred to as euthyroid Graves' disease
 - * Euthyroid is a term meaning that thyroid function tests are normal. Most people with euthyroid Graves' disease develop a thyroid disorder within eighteen months of the emergence of the eye disorder
 - * But some people with euthyroid Graves' disease never develop thyroid dysfunction

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Thyroid Eye Disease

- ~ What causes the Thyroid Eye Disease signs and symptoms?
- ~ The high and low levels of T3 and T4
- ~ The antibodies that are attacking the thyroid gland

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Thyroid Eye Disease

- ~ Thyroid Eye Disease has 2 phases
 - * A phase secondary to abnormal thyroid hormone levels
 - Increased or decreased FT3 and FT4 levels
 - Once these levels are normalized, ocular symptoms will resolve
 - * Congestive Autoimmune form of Thyroid Eye Disease
 - Active phase-stimulating or blocking TRAb are causing ocular activity
 - Plateau phase-reduced activity
 - Resolution phase-symptoms regress and eyes return to normal

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Phase secondary to abnormal thyroid hormone levels (T₃/T₄) (Thyroid Eye Disease)

<ul style="list-style-type: none"> ~ Hyperthyroidism eye symptoms <ul style="list-style-type: none"> * Excess hormone acting on the nerves that supply the eye * Usually spastic and include staring * Dryness * Eyelid retraction 	<ul style="list-style-type: none"> ~ Hypothyroidism eye symptoms <ul style="list-style-type: none"> * Deficient hormone causing venous congestion, impaired circulation and fluid stagnation * Periorbital edema
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- ~ This form of TED resolves within a few weeks after thyroid hormone levels (FT4 and FT3) are corrected and brought back into the normal range
- ~ The pituitary hormone TSH can stay low or suppressed for many months during the course of treatment for hyperthyroidism and doesn't mean that the patient is still hyperthyroid
- ~ TSH also lags at least 6 weeks behind thyroid hormone levels and often remains elevated longer in people who have been hypothyroid
- ~ Relying on the TSH level can be misleading and in treating TED

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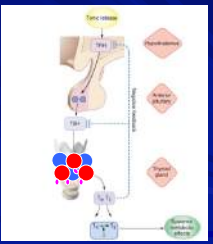
Congestive Autoimmune form of Thyroid Eye Disease (Active phase, Plateau phase, Resolution phase)

- ~ Caused by both stimulating and blocking TSH receptor antibodies (TRAb) and also immune system chemicals known as cytokines
- ~ Secondary targets appear to be TSH receptor antigens (epitopes) located on orbital fibroblasts as well as dermal fibroblasts
- ~ Active "inflammatory" phase of TED varies
 - * Symptoms resolve quickly although on average the active phase lasts about 12-18 months
 - * TRAb levels are high, patients are smokers, nutrient deficiencies are present, or the patient continues to be exposed to environmental triggers such as excess dietary iodine, the active phase can last as long as 5 years
 - * Avoid any lid, muscle or orbital surgery
- ~ Plateau phase and Resolution "Passive" phase
 - * An individual may be left with structural changes, such as eye protrusion, eyelid retraction, and in some cases, double vision
 - * There are corrective procedures that can be performed to address these problems

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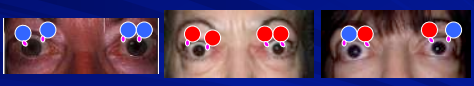
Euthyroid Graves' disease

~ If thyroid function is normal. How does one develop thyroid eye disease?

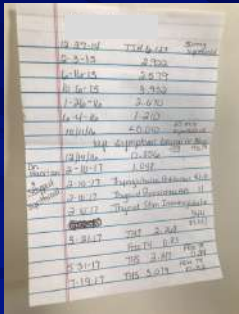


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Similar receptors are found in the skin, fat and muscle of the orbit



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You're in the Know

Normal Values
Thyroglobulin 20 IU/ml
Peroxidase <35 IU/ml
TSH 1.75 IU/ml

It does work!

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General Ocular Symptoms

- ~ Prominent eyes, stare
- ~ Pain
- ~ Lacrimation
- ~ Eyelid swelling
- ~ Foreign-body sensation
- ~ Double vision
- ~ Photophobia
- ~ Decreased vision in one or both eyes

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NOSPECS: Grading System

- ~ 1969 by S.C. Werner
- * Class 0: No signs or symptoms
- * Class 1: Only signs, upper lid retraction
- * Class 2: Soft Tissue Involvement with symptoms
- * Class 3: Proptosis
- * Class 4: EOM Involvement
- * Class 5: Corneal Involvement
- * Class 6: Sight Loss

~ Class 2-6 document severity

- * 0: absent
- * A: minimal
- * B: moderate
- * C: marked

~ Within classes 2 to 6 the investigator has to differentiate the severity grades 0, A, B, C

~ NOSPECS, classifies severity but not the activity or stage (active/inflammatory or passive/congestive)

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NOSPECS: Grading System

- ~ 0: No symptoms or signs
- ~ 1: Only signs (upper lid retraction without lid lag or proptosis)
- ~ 2: Soft tissue involvement with symptoms (excess lacrimation, sandy sensation, retrobulbar discomfort)
 - * Grade 0: absent
 - * Grade A: minimal (edema of lids, injection, sandy feeling)
 - * Grade B: moderate (edema of lids, injection, chemosis, FBS, pain behind eyes)
 - * Grade C: marked
- ~ 3: Proptosis associated with classes 2-6 only
 - * Grade 0: absent
 - * Grade A: minimal: 21mm -23mm
 - * Grade B: moderate: 24mm -27mm
 - * Grade C: marked: 28mm or more
 - * Specify if inequality of ≥ 3 mm between eyes, or if progression of ≥ 3 mm under observation

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NOSPECS: Grading System

- ~ 4: EOM involvement (usually with diplopia)
 - * 0: absent
 - * A: minimal (limitation of motion, patient reports diplopia but no obvious restriction)
 - * B: moderate (evident restriction of motion)
 - * C: marked (position of globe is fixed)
- ~ 5: Corneal involvement (due to proptosis, incomplete closure, lagophthalmos)
 - * 0: absent
 - * a: minimal (staining)
 - * b: moderate (ulceration)
 - * c: marked (clouding, necrosis, perforation)
- ~ 6: Sight loss (due to optic nerve involvement)
 - * 0: absent
 - * A: minimal (disc pallor or edema, or VF defect, vision 20/20-20/60)
 - * B: moderate (same as A but VA 20/70-20/200)
 - * C: marked (blindness, VA < 20/200)

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LEMO Classification

- ~ 1991-Boergen and Pickardt
- ~ Complements NOSPECS
- ~ 4 finding-categories
 - * Lid
 - * Exophthalmos
 - * Muscular
 - * Optic nerve
- ~ Grade between 0 and 4 depending on severity
- ~ LEMO, classifies severity but not the activity or stage (active/inflammatory or passive/congestive)

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LEMO Classification

Lid (L)

- 0: missing
- 1: lid edema only
- 2: real retraction (Impaired lid closing)
- 3: retraction and upper lid edema
- 4: retraction and global lid edema

Exophthalmos (E)

- 0: missing
- 1: eye closing not impaired
- 2: conjunctival injection in the morning
- 3: persistent conjunctival injection
- 4: corneal complications

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LEMO Classification

Muscular (M)

- 0: missing
- 1: detectable in imaging only
- 2: Pseudoparesis
- 3: Pseudoparalysis

Optic Nerve (O)

- 0: missing
- 1: regarding color vision only or detected via VEP
- 2: peripheral scotoma
- 3: central scotoma

L1EIM200
Endocrine ophthalmopathy with lid edema, exophthalmos, pseudoparesis of external eye muscles, and no optic nerve involvement

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Clinical Activity Score (CAS)

Thyroid disease characterized by:

- * Severity
- * Activity – want 3 or above
- CAS (1-7)

Studies by Tepezza

Payers using CAS for approval

- * Due to wide open label
- * Those infusing are charting the CAS

Clinical Activity Score	
1	Painful feeling behind globe
2	Pain on retrobulbar globe
3	Redness of eyelids
4	Redness of conjunctiva
5	Chemosis
6	Inflammatory eyelid swelling
7	Inflammation of conjunctival sac
8	Increase of 2 mm in proptosis in last 2-3 months
9	Decrease in visual acuity in last 3-7 months
10	Decrease in eye movements of 30° in last 1-3 months

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Lid Involvement

- ~ Lid Retraction
- ~ Lid Lag
- ~ Lagophthalmus

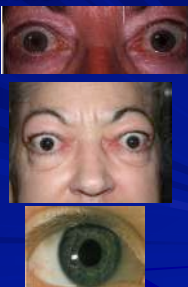
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Lid Retraction

- ~ Scleral show in primary gaze
- ~ Most commonly seen complication
- ~ Occurs in ~90% of Grave's patients
- * Excess stimulation of Muller's muscle
- * Fibrotic inferior rectus
- * Mechanical restriction or infiltration of levator
- * Increased orbital volume causes exophthalmos

Normal Lid Position

- * Upper lid intersects cornea at the 2 and 10 o'clock positions
- ~2 mm below the limbus
- * Lower lid coincident or 1-2mm below the limbus



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Eyelid Lag: von Graefe's Sign

- ~ Immobility or lagging of upper eyelid on downward gaze
- ~ Fibrosis of the inferior rectus muscle may induce lower lid retraction



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Lagophthalmos

- ~ Inability to form a complete lid closure with a normal blink due to Exophthalmos/ Proptosis
- ~ Often leads to corneal exposure

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Soft Tissue Involvement

- ~ Conjunctiva
- ~ Chemosis
- ~ Periorbital edema

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Conjunctiva

- ~ Conjunctival and episcleral injection
 - * Especially near the horizontal recti insertions
- ~ Chemosis
 - * Edema of the conjunctiva and caruncle
- ~ Superior Limbic Keratoconjunctivitis
 - * 65% correlation between SLK and systemic thyroid disease
 - * Rheumatoid arthritis
 - * Sjögren's syndrome



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"If it is Red think TED"

Dr. Andy Morgenstern 12-7-2013, OMS-Contemporary Resort



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Periorbital Edema

- ~ Inflammation of the subcutaneous connective tissue
- ~ May be first sign of thyroid eye disease
- ~ Greatest in the morning

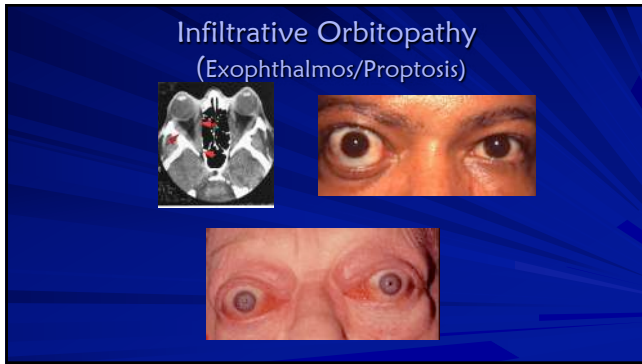


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Infiltrative Orbitopathy (Exophthalmos/Proptosis)

- ~ Thyroid Eye Disease is most common cause of unilateral and bilateral exophthalmos
- ~ The term exophthalmos is reserved for prominence of the eye secondary to thyroid disease
- ~ May need MRI to determine or obvious exophthalmos may be present
- ~ It is permanent in 70% of cases
- ~ Caused by increased volume of the extra ocular muscles
 - * Lymphocytic infiltration
 - * Proliferation of fibroblasts
 - * Edema within the interstitial tissue of the muscle

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Exophthalmometry

- ~ Is race dependent (Asians versus Black men is statistically significant)
- ~ Hertel or Luedde results
- ~ Adults
 - * Average reading 17 mm
 - * 95% of population have readings between 13-21mm
- ~ General concerns
 - * A difference of 2 mm or more between the eyes
 - * A measurement of more than 24 mm

Race	Mean Normal Value	Upper Limits
	mm	mm
White women	15.4	20.1
White men	16.5	21.7
Black women	17.8	23.1
Black men	18.5	24.7
Asians	----	18.0

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Restrictive Myopathy

- ~ Secondary to edema and fibrosis of EOM's
- ~ Inferior Rectus (IR) muscle is most commonly involved
- ~ Occurs in 30-50% of patients
- ~ Diplopia may be transient but in 50% it's permanent


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IOP in Thyroid Eye Disease

- ~ A rise in IOP has been reported with TED
- ~ I would have higher suspicion when you see
 - * Periorbital edema
 - * Exophthalmos, proptosis
 - * Restrictive myopathy
- ~ Some literature reports IOP in up gaze to be part of the diagnoses of thyroid dysfunction

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Restrictive Myopathy

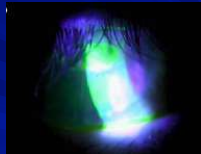


Obvious restrictive myopathy but also note the periorbital edema, and conjunctival hyperemia

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Corneal Exposure

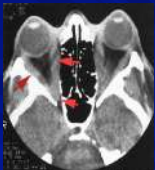

Exposure keratopathy secondary to exophthalmos and lagophthalmos
Significant threat to visual function



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Optic Neuropathy

Affects 5% of patients
Usually mild to moderate exophthalmos and shallow orbits
Enlargement of the recti muscles compresses ONH or its blood supply at the apex of the orbit
Compression MAY occur without significant proptosis
Compressive and/or ischemic and/or toxic

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
Treatment of Thyroid Eye Disease

Depends on what phase of the disease we are in:
Phase secondary to abnormal thyroid hormone levels
Active "inflammatory" phase
Plateau phase and Resolution "Passive" phase
Depends on what orbital tissue or structures are involved
Depends on the risk of vision loss
Depends if primary, secondary or tertiary thyroid dysfunction
Management consists of:
Control of inflammation
Prevention of ocular and visual damage
Addressing ocular motor abnormalities
Improving cosmetic disfigurement
Patient education is essential
Communication with an endocrinologist or internist will ensure proper patient care

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Treatment of Thyroid Eye Disease

Palliative (hormone imbalance, active, passive)
Lubricants
Topical anti-inflammatory (Lotemax/Restasis)
Prisms
Steroids (active phase)
Orals
Peri-ocular injections
IV with oral steroid taper
Orbital radiotherapy (active phase)
Orbital Decompression (passive phase)
Fat removal orbital decompression (FROD)
Large orbits
Bone removal orbital decompression (BROD)
Small orbits
Both FROD and BROD



Smoking causes the thyroid eye disease to be more severe
Smoking causes treatments to be less effective

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Treatment of Thyroid Eye Disease

Paradigm shifts
Decrease in orbital radiotherapy
Waiting for passive stage but doing surgery
Increase usage of fat removal orbital decompression as first approach
Peri-orbital injection of steroids for recurrent disease after orals
Future
Looking for better or different ways to treat the active phase of this disease

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Lid Retraction, Eyelid Lag, Lagophthalmos

- ⚡ Must treat underlying thyroid dysfunction
- ⚡ Abnormal hormone level and Active phase
 - ★ Treat the exposure keratitis with lubricants
 - ★ Tape eyelids shut at night
 - ★ Lid weight
 - ★ Moisture chamber at night
 - ★ Antibiotic ointments
- ⚡ Passive Phase
 - ★ Surgical Management
 - ★ Inferior rectus recession
 - ★ Mullerotomy
 - ★ Recession of lower lid retractors



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Lid Retractor Surgery



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Conjunctiva, Periorbital edema


- ⚡ Topical lubricants
 - ★ Artificial tears
 - ★ Ointments at night
 - ★ Topical steroids
 - ★ Restasis?
- ⚡ Tape eyelids closed at night or use mask
- ⚡ Elevate head at night to decrease lid edema
- ⚡ Oral diuretics Acetazolamide
- ⚡ Oral steroids
 - ★ 60-80mg/day for 3 months
- ⚡ IV steroids
- ⚡ Periorbital steroids
 - ★ Kenalog last 1 month



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Infiltrative Orbitopathy (Exophthalmos/Proptosis)


- ⚡ Orbital Disease Consult
 - ★ Systemic steroids to reduce inflammation
 - ★ Low dose radiotherapy
 - ★ Surgical orbital decompression



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Restrictive Myopathy

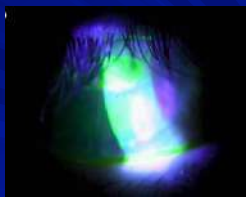
- ⚡ Non-surgical (while waiting for stability)
 - ★ Teach proper head position to alleviate diplopia
 - ★ Prism in spectacle correction (Fresnel or ground in)
 - ★ Oral steroids
 - ★ Botulinum toxin injection
- ⚡ Surgical Consult
 - ★ Recession of the rectus muscle/s involved
 - ★ Diplopia in primary gaze, reading gaze or both
 - ★ Stable angle of deviation for at least 6 months
 - ★ No evidence of active disease
 - ★ Binocular vision in at least primary and reading positions



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Corneal Exposure



- ⚡ Manage the corneal defect as first line
 - ★ Lubricating and antibiotic
 - ★ Lid taping
 - ★ Moisture barrier
- ⚡ Orbital Disease Consult
 - ★ High dose oral steroids
 - 120-140mg /day x 7 days
 - ★ Orbital decompression



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Optic Neuropathy

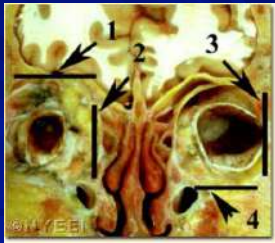
- ~ Systemic Steroids
 - * If rapidly progressive and painful in the early stage of the disease
 - * Only if no contraindications
 - * Prednisolone 80-100mg, expect results within 48hrs. Taper dose and d/c within 3 mo
- ~ IV Methylprednisolone
- ~ Radiotherapy; if contraindication to steroid
- ~ Orbital decompression

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Orbital Decompression

- ~ Not effective if no medical treatment
 - * Two-wall decompression
 - 3-6 mm retro-placement of the globe
 - * Three-wall decompression
 - 6-10mm retro-placement
 - * Four-wall decompression
 - 10-16mm retro-placement



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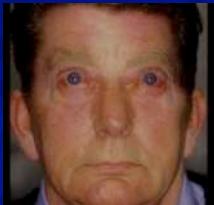
Orbital Decompression (Surgical/Cosmetic)



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

Thyroid Eye Disease and Depression

~ When facial disfigurement occurs, thyroid eye disease is equivalent to the diagnosis of cancer and AIDS



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Orbital Decompression (Medical/Vision Threatened)

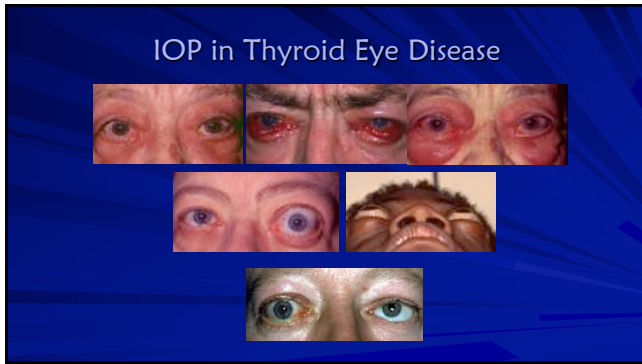



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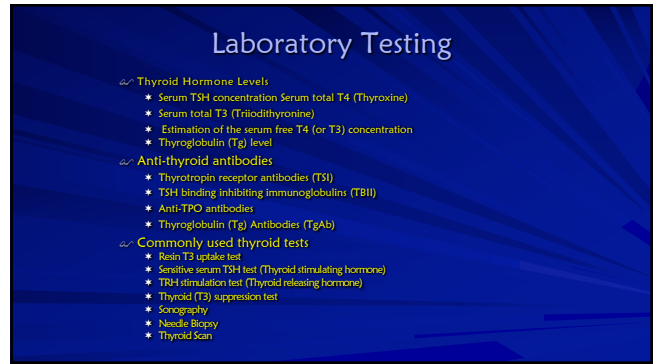
IOP in Thyroid Eye Disease

- ~ A rise in IOP has been reported with TED
- ~ I would have higher suspicion when you see
 - * Periorbital edema
 - * Exophthalmos, proptosis
 - * Restrictive myopathy
- ~ Some literature reports IOP in up gaze to be part of the diagnoses of thyroid dysfunction....let's discuss

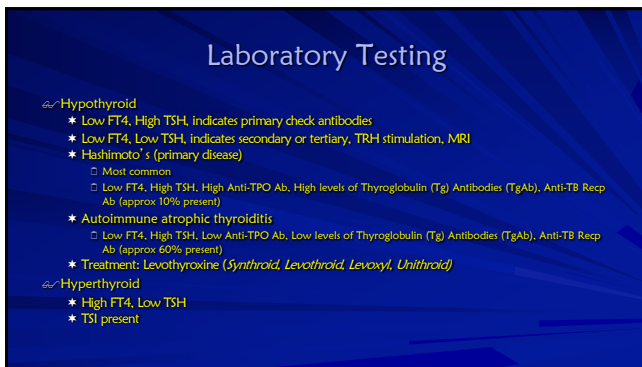
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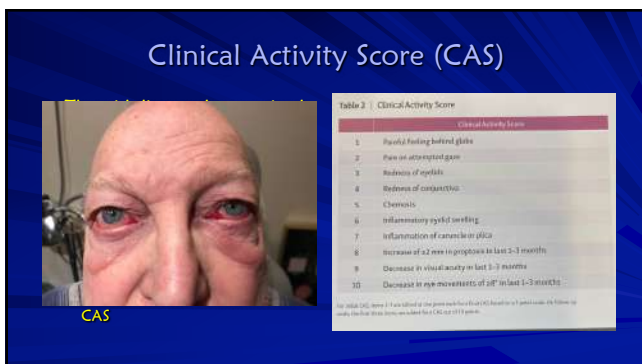
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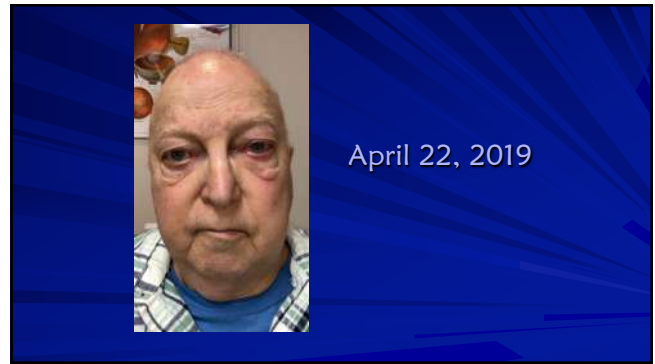
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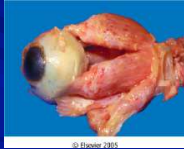
100



101

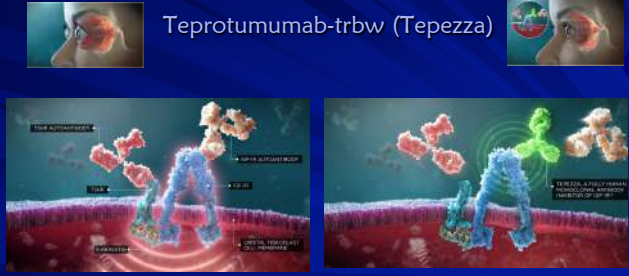
Teprotumumab-trbw (Tepezza)

- ~ Horizon Therapeutics – HQ Dublin, Ireland and US based Chicago
- ~ Biologic pharmaceutical
 - * Chinese Hamster Ovary
 - * Infusion, 8 total, every 3 weeks
- ~ Thyroid eye disease
 - * IGF1 (Insulin like growth factor 1) and TSH receptors are over expressed
 - * IGF-1 receptor inhibitor monoclonal antibody
 - o On the orbital fibroblasts
 - Inhibiting downstream inflammatory cascade
 - Cytokines, hyaluron, leukotriene
 - Differentiation into adipocytes and myofibroblasts
- ~ Phase 2 and published in New England Journal of Medicine
- ~ Phase 3 completed
 - * Published - New England Journal of Medicine
- ~ PDUFA- March 2020, was approved early in 2020



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Teprotumumab-trbw (Tepezza)




<https://www.tepezza.com/hcp/tepezza-mia/>

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Immunosuppression?

~ **Biologics**

- * **Immunosuppression biologics** – suppress the immune system to get the effect
 - Remicade – “1st generation”
 - Chimeric molecule – mouse and human protein, a lot of sensitivity
 - Humira
 - Anti-TNF (RA and Crohn’s Disease)
 - Fully human protein, less sensitivity
 - Rituxan
 - CD 20 suppressor (B cell suppression)
 - Actively suppress the immune system
- * **Immunomodulatory**
 - Tepezza
 - IGF-1R inhibitor
 - Full humanized monoclonal antibody
 - All the proteins are human – less to no sensitivity – more focused effect
 - Orbital fibroblasts to myofibroblast or adipocytes
 - Hyaluronic acid, glycosaminoglycan



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Teprotumumab-trbw (Tepezza)

~ **Optics and Optic-X Studies**

- * 8 infusions, every 3 weeks, 24 weeks
- * Optics – acute, less than 9 months of disease
- * Optics X – chronic, 12-16 months disease

~ **Clinical Activity Score**

- * Spontaneous pain, gaze evoked pain, eyelid erythema, chemosis, inflammation
- * Scale of 7, needed 4 to be in the study

~ **Proptosis**

- * Improvement of 2 mm or better

~ **Diplopia**

- * Scale of 0, 1, 2, or 3

~ **Grave’s Ophthalmopathy -Quality of Life Score**

- * Scale 0-100

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Teprotumumab-trbw (Tepezza)

~ **Clinical Activity Score (CAS)**

- * Spontaneous pain, gaze evoked pain, eyelid erythema, chemosis, inflammation
- * Scale of 7, needed 4 to be in the study
 - 78% improved to 0 or 1, 7% improved 0 or 1 with placebo

~ **Proptosis**

- * Improvement of 2 mm or better
 - 83% had 2 mm or better, 10% with placebo
 - Average was 3.2 mm at week 24

~ **Diplopia**

- * Scale of 0, 1, 2, or 3
 - 68% improved 1 point, 29% with placebo

~ **Grave’s Ophthalmopathy -Quality of Life Score**

- * Scale 0-100
 - 17.28 point improved, 1.80 with placebo

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Teprotumumab-trbw (Tepezza)

~ **Adverse Reactions**

- * **Very well tolerated**

* The most common adverse reactions (incidence ≥5% and greater than placebo) are muscle spasm, nausea, alopecia, diarrhea, fatigue, hyperglycemia, hearing impairment, dysgeusia, headache, and dry skin.

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Teprotumumab-trbw (Tepezza)

~ **Infusion Reactions (mild/moderate):** approximately 4% of patients

- * transient increases in blood pressure, feeling hot, tachycardia, dyspnea, headache, and muscular pain
- * consideration should be given to premedicating with an antihistamine, antipyretic, or corticosteroid and/or administering at a slower infusion rate.

~ **Hyperglycemia:** Increased blood glucose or hyperglycemia

- * In clinical trials, 10% of patients experienced hyperglycemia
- * Monitor patients for elevated blood glucose and symptoms of hyperglycemia while on treatment with teprotumumab
- * Patients with preexisting diabetes should be euglycemic before beginning treatment

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Teprotumumab-trbw (Tepezza)

~ **Infusion center**

- * Go to Horizon website
- * Contact Us
- * Type in your question
 - Looking for infusion center

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Biologics Used Off Label for TED

Small Molecule Therapy	Target	Dosing	Outcomes	Side Effects
Humira®	CD20	2 infusions of 400 mg each, 2 weeks apart	Most result in improvement of CAS, ptosis, and orbitopathy	Localizations of inflammatory based disease, infections, hepatomegaly
Abatacept®	TNF-α	Subcutaneous injection of 100 mg every 2 weeks for 48 weeks	Most showed decrease in inflammation or disease progression or both in the study	Upper ODS
Infliximab	TNF-α	Infusions at 5 mg/kg each dose over 2 years	Case reports of disease improvement in small studies. CAS, ODS, and orbital complications resolved in 1 case after 2 doses	Infections, malignancies especially lymphoma, drug-induced liver
Tocilizumab	IL-6	3 infusions at 8 mg/kg given every 4 weeks	ORX with 30 mg/kg treatment in CAS, most studies resolution of 25% or more of ODS	High incidence of liver, hematologic, pulmonary
Trastuzumab	HER-2	Initial infusion of 12 mg/kg, followed by 6 infusions at 6 mg/kg given every 3 weeks	Reduced symptoms in 78% of patients, improved CAS, ODS, and CAS. ODS improved in 83%	Most common: acute myeloid leukemia, MDS, diarrhea, lung disease, febrile neutropenia, and fatigue. However, 25-50% of patients require stopping study or treatment

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Key Tenants of Aging, Performance and Vitality

- 🔥 Oxidative Stress / Inflammation
- ⚖️ Hormonal Balance
- 🌞 Stress Hormones
- 📊 Glucose / Insulin Regulation
- 🌱 GUT integrity and microbiome diversity
- 🛡️ Immune Balance
- 🌍 Environmental Exposure/Burden
- 🌟 Individuality

Credit to: James LaValle, RPh, CCN

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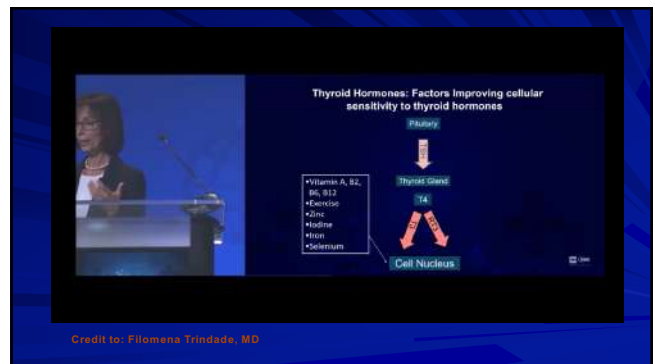
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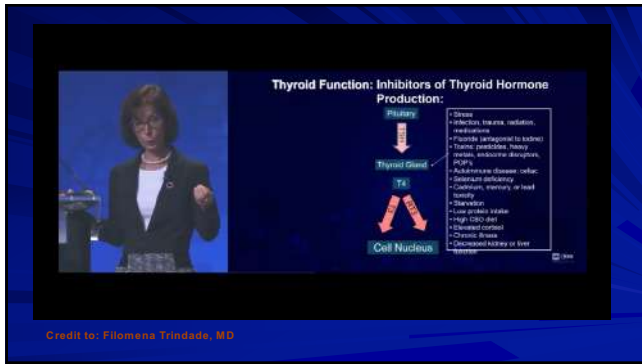
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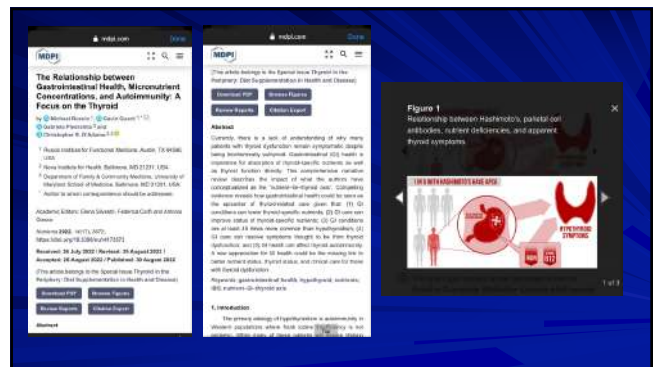
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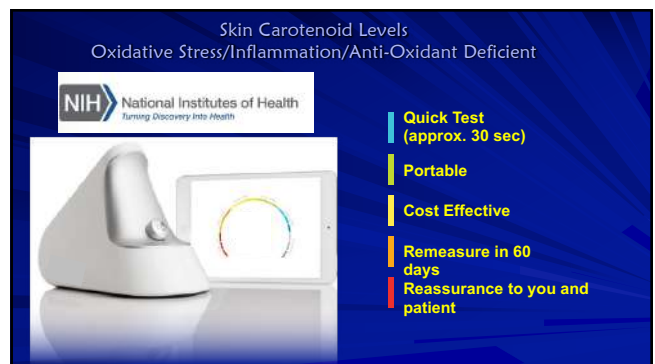
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Ingredients			Amount			% Daily Value		
Ingredients								
Non-Proprietary								
Shelf-Stable TPACKET								
Vitamin A (BPA) as Beta Carotene (BPA) from Marisbio (BPA) and Vitamin A palmitate (BPA) (BPA)	2750 mg (BPA)	250%						
Vitamin C (as Cellulose Ascorbate)	2100 mg	221%						
Vitamin D (as Cholecalciferol)	2.000 µg	25%						
Vitamin E (as Hydroxytolerol)	10.3 mg	19%						
Vitamin K (as Phytylphosphonate)	21 mg	2%						
Zinc (as Zinc Oxide Monohydrate)	2.5 mg	15%						
Biotin (as Biotin)	4.25 mg	22%						
Niacin (as Nicotinamide)	175 mg	300%						
Vitamin B6 (as Pyridoxine Hydrochloride)	5 mg	104%						
Vitamin B12 (as Cyanocobalamin)	1000 µg	20%						
Iron (as Ferrous Fumarate)	25 mg	250%						
Calcium (as Calcium Carbonate, Di-Calcium Malate, Calcium Ascorbate)	300 mg	30%						

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Supplement Facts		
Supplement Facts		
Serving Size: 1 Tablet		
Amount Per Serving		
		% Daily Value
Vitamin A (as Retinyl Palmitate)	10,000 IU	200%
Vitamin B12 (as Methylcobalamin)	1,000 µg	200%
Vitamin C (as Ascorbic Acid)	2,000 mg	400%
Vitamin D (as Cholecalciferol)	25 µg	500%
Vitamin E (as d-Alpha-Tocopherol)	100 mg	200%
Vitamin K (as Menaquinone-2)	100 µg	200%
Zinc (as Zinc Oxide)	25 mg	500%
Iron (as Ferrous Fumarate)	25 mg	500%
Biotin (as D-Biotin)	5,000 µg	100,000%
Niacin (as Nicotinamide)	175 mg	350%
Cyanocobalamin	1,000 µg	20,000%
Calcium (as Calcium Carbonate)	300 mg	30%
Calcium (as Calcium Malate)	0 mg	0%
Calcium (as Calcium Ascorbate)	0 mg	0%

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Key Tenants of Aging, Performance and Vitality

- Oxidative Stress / Inflammation
- Hormonal Balance
- Stress Hormones
- Glucose / Insulin Regulation
- GUT Integrity and microbiome diversity
- Immune Balance
- Environmental Exposure/Burden
- Individuality

Credit to: James LaValle, RPh, CCN

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Signs in Thyroid Eye Disease

- Dalrymple's sign: Lid retraction
- von Graefe's sign: Upper lid lag on downward gaze
- Griffith's sign: Lower lid lag on downward gaze
- Boston's sign: Jerky irregular movement of the upper lid on downward gaze
- Jellinek's sign: Increased pigmentation of the lids
- Stellweg's sign: Infrequent blinking
- Kocher's sign: Increased lid retraction with visual fixation
- Enroth's sign: Puffy swelling of the lids
- Rosenbach's sign: Tremor of closed lids
- Mobius' sign: Weakness of convergence
- Ballet's sign: Palsy of one or more extraocular muscles
- Suker's sign: Weakness of fixation on lateral gaze
- Cowen's sign: Jerky papillary contraction to consensual light
- Knies' sign: Unequal dilatation of the pupils
- Jeffrey's sign: Absence of forehead wrinkling on upward gaze

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Questions and Thank You!

**Thyroid and Thyroid Eye Disease
Clinical Pearls and Innovations for 2023**

Greg Caldwell, OD, FAAO
Phoenix, AZ
Saturday, April 15, 2023

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