

1


3

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2


## Diagnostics



## Clinical diagnosis

PE and symptoms as clues
General diagnostics
Infectious triggers
Environmental triggers

## Neuro P/E relevant to BGE

Burdened appearance
Dilated pupils
Hypotonia
Motor apraxia
Dyspraxia
Normal strength
Normal reflexes, not hyperactive as in Wilson's dz
Abnormal movements
Chorea
Choreiform movements, not age appropriate
Tics
Steriotypies
Ballismus
Overall rational irrationality (they realize or have insight into the abnormality)
Presented by Dr. Elizabeth Latimer
Autoimmune Encephalitis Post-Streptococcal Evaluation \& Treatment Conference Oct 2019



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## Diagnostics



Clinical diagnosis
PE and symptoms as clues

General diagnostics
Infectious triggers

Environmental triggers



13

## Autoantibodies

Antinuclear antibodies multiplex, reflex to dsDNA, RNP, Sm, SS-A, SS-B
Demyelination Antigens
Anti-tubulin $\lg M / \lg G+\lg A$
Anti-myelin basic protein $\lg \mathrm{M} / \mathrm{IgG}+\lg A$
BBB Disruption ~
Anti s100b $\operatorname{lgM} / \operatorname{lgG}+\operatorname{lgA}$ (*increases with exercise)
Optical and ANS Disorders ~
Anti-neuron specific enolase $\lg M / \lg G+\lg A$
Peripheral Neuropathy -
Anti-GM1 lgM/IgG+lgA
Anti-GM2 $\operatorname{lgM} / / \lg G+\lg A$
Brain Autoimmunity ~
Anti-HSV1 IgM/gG+IgA
Anti-cerebellum IgM/gG+IgA
Anti-purkinje cell lgM/IgG+lgA
Anti-pituitary antibodies (APA) (hypophysitis post TBl)

## Immunocompetence

Quantiative lgGAME with lgG subclasses
Red top tube or SST? Depends on goals for testing
Serum separator will bind some antibodies and under-report, esp in those with low Ig's.
(4. Dr. Paul Anderson)

NOTE that all antibody-based testing will be affected by IVIG, including other autoimmune and infection
Lymphocyte Subset/Differential Panel ~
Offers the advantage of detecting the cell type that causes the immune defect.
3 types of lymphocytes: B, T and NK cells.
All share the same progenitor cells: hematopoietic stem cells in the bone marrow, which then give rise to multipotent progenitors, to early lymphocyte progenitors (ELP) and eventually to the differentiated
progenitors of NK, B or T cells. $B$ and

NK cells, in contrast, are not antigen-specific lymphocytes, thus belonging
to the innate immune system.
PMO: 3 2028274



|  | Table 2 Lymphocyte subset percentages in healthy children: Distribution by age (reproduced with permission from Shearer et al ${ }^{18}$ ) |  |  |  |  |  |  |  |  | $\sim^{20238 \text { orilic Cista }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Subset | $n$ | $\begin{gathered} 0-3 \text { months } \\ \begin{array}{c} \text { Median } \\ \text { (range) } \end{array} \\ \hline \end{gathered}$ | $\begin{gathered} 3-6 \text { months } \\ \text { Median } \\ \text { (range) } \\ \hline \end{gathered}$ | $\begin{gathered} \text { 6-12 months } \\ \text { Median } \\ \text { (range) } \\ \hline \end{gathered}$ | $\begin{aligned} & 1-2 \text { years } \\ & \text { Median } \\ & \text { (range) } \\ & \hline \end{aligned}$ | $\begin{aligned} & 2-6 \text { years } \\ & \text { Median } \\ & \text { (range) } \end{aligned}$ | $\begin{gathered} \begin{array}{c} \text { c-12 years } \\ \text { Mediar } \\ \text { (range) } \end{array} \\ \hline \end{gathered}$ | $\begin{gathered} \text { 12-18 years } \\ \text { Median } \\ \text { (range) } \\ \hline \end{gathered}$ |  |
|  | 3 | 709 | 73 (53-84) | 66 (51-77) | 65 (49-76) | 65 (53-75) | 66 (56-75) | 69 (60-76) | 73 (56-84) |  |
|  | 19 | 709 | 15 (06-32) | 25 (11-41) | 24 (14-37) | 25 (16-35) | 21 (14-33) | 18 (13-27) | 14 (06-23) |  |
|  | $16 / 56$ | 784 | 8 (04-18) | 6 (03-14) | 7 (03-15) | 7 (03-15) | 9 (04-17) | 9 (04-17) | 9 (03-22) |  |
|  | 4 | 709 | 52(35-64) | 46 (35-56) | 46 (31-56) | 41 (32-51) | 38 (28-47) | 37 (31-47) | ${ }^{41}(31-52)$ |  |
|  |  | 709 | 18 (12-28) | 16 (12-23) | 17 (12-24) | 20 (14-30) | 23 (16-30) | 25 (18-35) | 26 (18-35) |  |
|  | $4 / 45 \mathrm{RA} /$ <br> 62 L | 805 | 89 (61-94) | 88 (64-92) | 83 (58-91) | 79 (62-90) | 70 (50-85) | 58 (42-74) | 51 (31-65) |  |
|  | ${ }_{8 / 45 R A}$ <br> 62 L | 807 | $79(56-88)$ | 77 (53-88) | 72 (47-87) | 71 (46-85) | 64 (42-81) | 58 (39-73) | 56 (42-73) |  |
|  | 4/45RA | 805 | 90 (64-95) | 90 (77-94) | 86 (64.93) | 81 (63-91) | 71 (53-86) | 59 (46-77) | 53 (33-66) |  |
|  | 845RA | 887 | ${ }^{93}(80-99)$ | $94(85-98)$ | 91 (75-97) | 88 | ${ }^{86}$ (69-97) | $80(63-92)$ |  |  |
|  | 4/DR/38 | ${ }_{8}^{805}$ | 3(01-06) | 4 (02-09) | 4 (01-09) | 5(02-09) | 5 (02-09) | $4(01-08)$ | 3(02-06) |  |
|  | 8/DR/38 | 807 | 5 (02-17) | 7 (03-16) | ${ }_{5}^{8(03-25)}$ | 15 (05-30) | 13 (05-29) | $9(02-20)$ | 7 (03-18) |  |
|  | 4/DR | 805 | ${ }_{5}^{3(02-06)}$ | $5(02-10)$ | 5 (02-11) | 6 (02-11) | 7 (03-12) | $6^{6(03-13)}$ | 7 7(04-11) |  |
|  | 8/DR | 807 | 5 (02-20) | 7 (03-17) | 10 (04-27) | 16 (06-33) | 16 (07-37) | $12(06-29)$ | 12 (05-25) |  |
|  | $4 / 38$ | 805 | 98 (95-99) | 96 (90-98) | 95 (89-97) | $93(85-97)$ | 87 (74-94) | 79 (64-86) | 69 (50-79) |  |
|  | 8138 | 807 | 97 (89-99) | 95 (83-98) | 93 (78-98) | 91 (73-97) | 82 (52-93) | 70 (42-86) | $64(33-80)$ |  |
|  | ${ }_{8}^{428}$ | 806 | ${ }^{99} 9(95-100)$ | 99 (88-100) | 88 (90-100) | ${ }^{98}(94-100)$ | 98(92-99) | ${ }^{98}$ (92-100) | 97 (89-100) |  |
|  | $8 / 28$ $4 / 95$ | 806 806 | $76(54-87)$ $11(05-21)$ | $75(43-87)$ $14(08-21)$ | $70(42-83)$ $18(11-34)$ | - 63 ( 3 (11-31) | - $\begin{aligned} & 631(22-79) \\ & 31\end{aligned}$ | - $\begin{aligned} & 60(42-78) \\ & 39\end{aligned}$ |  |  |
|  | 8895 | 806 | 12 (02-33) | 15 (06-36) | 22 (08-47) | $31(07-50)$ | $34(12-57)$ | 33 (10-62) | 44 (15-71) |  |
|  | 3/4445RO | 676 | 10 (02-22) | 8 (03-16) | 9 (05-18) | 12 (07-20) | 16 (09-26) | 21 (13-30) | 28 (18-38) |  |
|  | 3/4-744RO | 672 | 3 (01-09) | $3(01-07)$ | 3 (01-08) | 6 (02-12) | $9{ }^{(04-16)}$ | $12(0421)$ | 13 (04-23) |  |
|  | 3/45RO | 676 | 14(03-31) | 13 (04-24) | $12(06-25)$ | 19 (09-31) | 27 (15-41) | ${ }^{33}(20-40)$ | ${ }^{41}(24-57)$ |  |
|  | $3-1 / 1988$ $3 / 119$ | 686 686 | - $50(13-75)$ | $66(00-82)$ $69(00-84)$ | $\begin{aligned} & 66(01-78) \\ & 67(01-80) \end{aligned}$ | $60(00-79)$ $63(00-80)$ | $\begin{aligned} & 55(01-70) \\ & 61(02-76) \end{aligned}$ | $39(00.60)$ $46(00-67)$ | $19(00-57)$ 21 (00-60) |  |
|  |  |  |  |  |  |  |  |  |  |  |
| CRISTA | PMD: 302482 |  |  |  |  |  |  |  |  |  |

 Table 2 Lymphocyte subset percentages in healthy children: Distribution by age (reproduced with permission from Shearer et al ${ }^{18}$ )

## Neuroquant MRI

Specialized MRI must be run at specific Neuroquant centers.

- TBAR with asymmetry
- may need to order Brain Development report b/c TBAR changes may reflect
neuronal development. (Dr. Gazda)
Does not require contrast. Age- and gender-matched controls.
Normal $=40-60$ percentile
May display enlargement of the caudate.
- Blue = edema/inflammation
- Red = atrophy

Thalamus >90\% mold and Lyme. (Dr. Ackerly)
Not ideal for child with tics, as they can't remain still for imaging.
Also not ideal for sound sensitive child or child who cannot tolerate ears being
covered.
Braces/retainers will alter findings.


## Food sensitivities

Proteins vs Peptides
Proteins ~
Measure immune system reactivity to whole, undigested, multi-dimensional (ie: 4D) proteins.
Challenges: only detect one aspect of "the elephant in the gut"
Limited to testing the water-soluble portions of proteins, leaving out non-water-soluble peptides (ie: gluten).
Peptides ~
Measure immune system reactivity to the small, typically not water-soluble, 2D peptides created when whole proteins are digested.
Reduces cross-reactivity; increased sensitivity because peptides are highly specific to the food from which they are derived.
Antibodies to a whole protein will not recognize or bind peptides, even if those peptides are found in that whole protein.
Clinically, testing for food sensitivities at the peptide level in addition to whole protein eliminates uncertainty around food reactions.


Neuroquant Triage Brain Atrophy Report TBAR


Genetic predispositions/expressions
PANS: HLA alleles:
HLA-B 38, 52, 55
My own observations:
Snps related to IgG: Fcy Receptors
Snps related to NTs: COMT, MAOA


Snps related to detox:
Phase I: CYP1A2, CYP1B1, CYP3A4 (mold)
Phase II: GSTM1, MTHFR, SUOX
Snps related to histamine: DAO
Metagenomics/metabolomics

## Drug metabolism genetics

Ultrasensitive to psychiatric medication ~
Due to BBB integrity or genetics?
Better to know child's drug clearance before prescribing, especially if the effect is slower metabolism and reduced drug clearance, concentrating the drug.
Testing provides Gene-Drug interaction chart.
Also be familiar with co-enzymes that up- or down-regulate that pathway. ie: B2, B6, NAD

## The fine art of insurance coverage

Set up for IVIG coverage in case it's needed in the future (analogous to starting an IV in the ER) **don't put PANDAS or PANS Dx in chart, unless you're in a state which mandates coverage** Test IgG and lgA (plus subclasses) ~

- Using SST tubes
- Test after 3 weeks without any integrative supportive measures (test the child's true nature.) - Test at the tail end of a steroid burst, if needed.

Also engage parent help. Bring child in every time they get sick to get it on the medical record.
Parent needs to keep school absence records, sports absence records, performance absences, etc. Cautions ~
Zinc lozenges, silver nasal sprays, propolis throat sprays turn positive Strep tests to negative. Diet, supplements, sleep routines, chiropractic adjustments, and all the other integrative treatments really work! We see in
the child, but bad for
A hiatus helps reveal the baseline.
Pneumococcal vaccine titers are not necessary, plus may be falsely lower in kids with
neumococcal vaccine titers are not necessary,
hypogam. Push back against insurance on this

## On the horizon?



## Metagenomics: NGS qPCR of brain/CSF

Metagenomic NGS is a novel diagnostic test with the potential to revolutionize the diagnosis of pediatric meningitis and encephalitis through unbiased detection of bacteria, viruses, parasites, and fungi in cerebrospinal fluid.
"We recommend NGS should be considered as a front-line diagnostic test in chronic and recurring presentations and, given current sample-to-result turn-around times, as second-line in acute cases of encephalitis."
PMID: 29305150, 34951470

## Group A Strep (GAS)

GAS is the dominant respiratory pathogen ~
Accounts for $20 \%-40 \%$ of cases of pharyngitis in children; the remaining are caused by viruses
GAS infections ~
Strep pharyngitis, otitis media, sinusitis, skin infections (perianitis)
Colonization posited in sinuses and Gl
GAS sequelae ~
Scarlet fever, cellulitis, necrotizing fasciitis, rheumatic fever, Streptococcal toxic shock syndrome, and post-Streptococcal glomerulonephritis

## Strep shoots the messenger

Rewires the immune system for its survival
Unique in its abilities to ~
Direct I/S remodeling in nose/throat (possibly perianally)
Promote its own replication
Alter I/S responses
Shoots the messenger": GAS virulence factors modulate maturation and survival of dendritic cells (DC) aka the "delivery" cells, effects that are likely to have a critical impact on activation of innate and adaptive immune responses

Only 6 of 24 GAS strains tested induced surface expression of MHC class II and costimulatory molecules consistent with DC maturation
The majority of the strains did not promote DC maturation and many triggered DC apoptosis.
pMID: 19712038
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## Perianal Strep Dermatitis (GAS)

Symptoms ~
Red rash around the anus with a well-defined margin
Sore rectum or anus
Anal pruritus
Pain with bowel movements or when wiping
Constipation
dx ~
Candidiasis, pinworms, eczema, and contact dermatitis from soaps, detergents,
and fragrances
Seasonality ~
Winter \& spring
Culture all perianal rashes AND culture to confirm successful treatment.
Not uncommon to have pharyngeal culture neg, but perianal positive.

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## Skin infections: Suppurative

Impetigo - honey-colored crust, superficial - heals without scarring.
Ecthyma - deeper lesion, below dermis, indolent. Starts as a pustule and erodes to an ulcer Often multiple lesions

Erysipelas - raised red rash with very sharp borders. In the lymphatics of the skin. Fever and pain from skin swelling. IV Abx.
Cellulitis - border vague and irregular. Skip areas/bare areas. Painful, may not have fever Associated with a break in skin.
Lymphangitis - rapidly progressive infection with initial cutaneous focus but spread of infection through lymphatics.

Necrotizing fasciitis/streptococcal myositis - Streptococcal gangrene. Superficial and possibly deep layers of muscles are killed. Pain and swelling are disproportionate to everything else. Needs surgery.
Streptococcal pupa fulminans - Skin and all structures underneath necrose. Blood vessels thrombose. + blood culture usually.



Strep "carriers" ~
Check parent/siblings tonsils
Often child's are small and parents/siblings are enlarged or boggy (may also be EBV)
Check skin infections of other family members
Pets ~
Animals cannot get infected by Strep as it's strictly a human pathogen
Transfer via saliva from licking carrier's face or skin infection
Probiotics ~
Until we know which peptide or protein induces an immune reaction, I recommend avoiding Strep-based probiotics

Neonatal (Group B)


Group B Streptococcus
Debated the degree to which Group B Strep in mom before birth contributed to the development of PANDAS, but doctors specializing in PANDAS have reported a correlation.

Vaginal swab culture

Why worry about Strep in PANS?


Strep is kryptonite in kids with PANDAS -andPANS
Even though it may not have been the triggering infection, Strep can trigger flares in PANS.

## Strep detection

Culture culture culture ~
Antibody response is more complex than previously understood. A negative rapid strep test can still be culture positive. F/U negative rapid with culture.
Rapid strep tests ~
Very high specificity ( $98-99 \%$ ) = very few false positives
However, sensitivity lower ( $90-95 \%$ ) = greater chance of false negatives.
Package insert recommends F/U negative tests with a culture.
Lawsuits against docs who didn't F/U with culture, missed Strep, and serious sequelae.
Cochrane Database Systematic Review 2016 ~
Out of 100 children with strep throat:
86 would be correctly detected with the rapid test
14 would be missed and not receive antibiotic treatment
s clinical over-reliance in rapid strep tests a contributing factor for the rise in PANDAS/PANS?
PMD: 27374000

## Mycoplasma pneumonia

Look for it, and look again, and again. It's ubiquitous and often asymptomatic or only mild illness.
Shares many of the same skills as Strep in evading the I/S and affecting the brain. Second most favorite places to play in the body are brain and CNS.

Can cause encephalitis. CNS complications are seen more so in kids. Just like Strep, certain proteins on Mycoplasma mimic brain tissue.

Been shown to be able to persist in an intracellular environment. Antibiotic resistance issues.
Do not develop lifelong immunity to Mycoplasma.
More common in the winter and is estimated to be much more common than previously understood.
A super-spreader: takes up to 3 weeks before symptoms develop and is shed from the respiratory tract for many weeks after symptom abatement.
Equates to up to 6 weeks of potential transmissibility with one infection.
Sometimes, Mycoplasma's main hideout is the tonsils. For children whose tonsils have become Mycoplasma reservoirs, removing the tonsils may be helpful.

## Mycoplasma pneumonia symptoms

Symptoms of respiratory Mycoplasma in children under 5
Watery eyes
Runny nose
Sore throat
Digestive changes such as diarrhea or vomiting
Symptoms of respiratory Mycoplasma in children 5 years or older:
Feel tired
Low-grade feve
Sore throat
May have a headach
Slowly worsening dry cough that may last for weeks
The cough is normally dry. Even though it commonly takes weeks for the cough to go away, it should stay dry. If the cough becomes productive, and is accompanied by worsening fever or chills, or feeling SOB, r/o "walking pneumonia".
Mycoplasma may cause other non-lung symptoms, such as achy muscles and joints, skin rashes, heart symptoms, iver inflammation, and eye symptoms such as pink eye and anterior uveitis.
Also mimics RBCs and can lead to hemolytic anemia. May be mistaken for Babesia which infects RBCs and causes many of the same circulatory symptoms.

## 45

## Chlamydia pneumonia

Respiratory infection, not the STI Chlamydia trachomatis,
Obligate intracellular bacteria that infects the respiratory epithelial tissue and may play a role in chronic inflammatory dzs.

Majority of individuals are exposed throughout their lifetimes with an antibody prevalence of $50 \%$ by age 20 and $80 \%$ by 60-70 years old.

Predominantly asymptomatic or mild, but can result in the development of acute upper and lower respiratory illness including bronchitis, pharyngitis, sinusitis, and pneumonia/community-acquired pneumonia.

CNS can also be a target.
Co-infection of $C$. pneumoniae and $M$. pneumoniae with SARS-CoV-2 is associated with more severe features
PMD: 30887565, 23218799, 11371700, 334882388

## Mycoplasma detection

IgG may or may not be positive with a positive IgM.
IgM remains positive much longer than other microbes, so can be a false positive.
Confirm IgM+ via immunofluorescence (Mayo) - titers vary wildly by the moment.
T-cell option


## C. pneumonia detection

T-cell
Chronic infection is somewhat more difficult to determine and requires the detection of persistent IgG levels, which is complicated by the fact that IgG has a half-life of weeks to months and may therefore be present for some time following acute infection.
It has been proposed that IgA levels may provide a better indication of chronic infection, but according to Dowell et al., the use of $\lg G$ and $A$ serological markers alone should not be used.
Identification of C . pneumoniae messenger RNA (mRNA) by PCR can also be used to determine whether C . pneumoniae is in a metabolically activated state.

PCR detects presence within tissues (ie: tonsils.)
PMD: 8865464, 11462188

## Bartonella spp

Include a number of different species, which are growing in number as detection improves.

Transmitted through flea bites, tick bites, and scratches or bites from an infected animal. Vertical transmission during pregnancy,
Best known as cat scratch fever, but that's misleading.
A scratch isn't required (vector bite),

- Not just cats - also dogs, rabbits, and many other pets such as hamsters and gerbils.

Can disperse far and wide in the body, and can migrate in and out of the blood. This causes a relapsing-remitting pattern, making it quite difficult to discern between a chronic Bartonella infection and a PANDAS or PANS flare.

Each species causes a slightly different symptom picture, which makes diagnosis difficult. There are commonalities based on favorite tissues to infect, those being the brain, nervous system, and connective tissue, especially collagen

## Bartonella

Formerly, considered an issue only for those with severe immune compromise.
New research is supporting what many of us working with P/P kids have found-it's far more common and often chronic. May even be "asymptomatic", but with the rising prevalence of anxiety and ypermobility in kids, one wonders about that.
Anxiety is the most common brain-related symptom with Bartonella, as well as neuropathies. We also see mood swings that can be quite drastic and seemingly unprovoked. These may present as out-of-control anger and rage events.
Regarding connective tissue, research suggests that Bartonella impairs collagen synthesis and repair. (So does mold.) This mechanism accounts for the "Bartonella stretch marks."

This collagen-interrupting effect is why many cases of hypermobility are due to undetected, chronic Bartonella in kids and teens living in moldy environments.
We worry about Bartonella's effect on the connective tissue of our hardest-working tissues, such as the eyes and the heart, where it can weaken and infect the heart valves.
Acute cases typically fit the classic $\mathrm{s} / \mathrm{sxs}$, but chronic Bartonellosis is missed frequently due to the variance in how it presents in different children.

## Acute Bartonella spp

Fever
ce-pick pains, especially of the ear
Skin rashes or nodules
Stretch marks that don't blanch
Extremely enlarged lymph nodes
Anger or rage events
Hemolytic anemia
Joint pain
Uveitis
Neuroretinitis
Encephalitis
Endocarditis
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Bartonella: Cutaneous lesion presentation depends on strain. "Bart striae" or non-blanching stretch marks. PMD: 33291688


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54


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## Bartonella diagnostics

Acceptable to treat based on a presumptive diagnosis
Famously difficult to detect due to their migratory pattern from the blood into tissues, evading the I/S.
FA, PCR, T-cell
"ILADS folklore" - draw between 2-4pm
May provoke with homeopathics for 1 week prior


57

## Prevention is Key

Attractors ~
$\mathrm{CO}_{2}$ is the tick attractor.
Also pheromones from Lyme carriers.
Certain mosquito-attracting (flavi)virus-induced skin volatiles:
Acetophenone, a volatile compound that is predominantly produced by the
skin microbiota, is enriched in the volatiles from the infected hosts to potently
stimulate mosquito olfaction for attractiveness.
An effect partially combatted by Vitamin A.
Defense ~
Treated clothing
Essential oil - lemon eucalyptus, yarrow (acaricidal); reapply often (min hourly)
Coming inside: clothes stripped and in hot dryer $\times 10 \mathrm{~min}$
Tape roll pets
Tick tubes around outdoor spaces
PMID: 35777 555, 38005473


58
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## "Never had a tick bite" "Not outdoorsy"

Tick saliva contains an anesthetic
Also anticoagulant and immunosuppressive substances
Provides a localized environment at the site of the bite to evade detection - enhancing infection

Soft-sided ticks (TBRF) are "snackers" - may self-detach and find new host, won't necessarily engorge

Migratory birds carry ticks anywhere the bird can go (even Home depot)


61


## Lyme: How To Make A Poultice To Extract

 Embedded Tick Head円ロ


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65


## Early disseminated

Onset weeks to months after bite.
Early disseminated Lyme can occur even if no acute sxs.
Areas ~
HT - carditis, A-V block
Neurological - cranial/peripheral neuropathy
M/S - migratory arthralgias
Eye - all the "itis"s - uveitis keynote, retinal tears
Skin/lymphatics
Liver/kidney - LFTs, proteinuria
Late or "chronic" Lyme
Onset months to years after tick bite.
Can also occur w/o any prior sxs.
Dr. Horowitz's Lyme/MSIDS Questionnaire is the premiere sx list.
May have never felt well since, or triggered by stressor (mold, MVA, surgery, dental, mental/
emotional, pregnancy, puberty, menopause, etc)
Correlated with extreme morbidity.
Chronic progressive multisystem illness in:
M/S
Neuro
Skin - acrodermatitis chronica atrophicans (European cisisa
Hormone

## Non-Lyme Borrelia - TBRF

Tick-borne relapsing fever
Transmission - soft-sided ticks (don't engorge,
they "snack" and may move hosts,) lice,
fleas, possible spider bites
Tests negative on Lyme disease tests.

In Children: Early TBRF

Before feeding
After feeding

Early Tickborne Relapsing Fever
High fever, chills
Headache
Muscle and joint aches
Fever relapses and lasts for about 3 days
Rarely a rash


70
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## In Children - Persistent/Chronic Borrelia

Persistent or chronic Lyme disease or Tickborne Relapsing Fever:
Fatigue
Brain fog
Problems remembering new learning
Child avoids play or friends
Mood changes, depression, anxiety
nsomnia
Headaches
requently changing vision
Rashes that come and go
Nerve pain, numbness, tingling, or random hot or cold feeling
Heart palpitations
Digestive problems
Muscle and/or joint pain that migrates around the body
requent musculoskeletal injuries
Generalized heightened body pain

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## The Great Imitators



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## Take-Aways

"yme" has become an umbrella term used to describe many iterations of tick-borne infections different infections/combinations of infections different stages/states
-tier reflex to WB missing an inordinate number of cases
ER/Urgent care labs too early to detect
Clinical diagnosis is sufficient to initiate Tx
Known tick bite is not required to Dx
Rash is not required to $D x$
Tx for $7-10$ days is not sufficient
Delayed onset of $T x$ is correlated to worse outcomes
Post treatment Lyme syndrome" is an erroneous Dx. IME culprit is surviving bacteria, but is blamed on the immune system gone awry
"Antibiotic refractory" - a research term - IME from undertreated and/or missed acute $\mathrm{Dz} \rightarrow$ widely disseminated, genetically savy bacteria (more later)
Reportable Dz - if they'll accept it (my story of Advanced Labs culture +)
Vertical transmission has been reported, positive cord blood and culture positive neonate


77

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Redien ${ }^{\text {PANDAS/PANS in the COVID-19 Age: Autoimmunity and }}$
Epstein-Barr Virus Reactivation as Trigger Agents?
Stefano Pallanti 12.2 and Michele Di Ponzio ${ }^{\text {sen }}$


mat $\stackrel{\circ}{\circ}$

Early antigen (EA) - add-on
Chronic/reactivated pattern ~ VCA-IgG - pos
VCA-lgM - neg
EA-IgG - pos
EBV-NA - highly pos
(if $3-4 \times$ positive, consider chronic/reactivated)
vs Past infection pattern ~
EA - neg
NA - lower pos


78
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## Influenza

Very commonly reported cause of PANS and flares by parents.
Influenza symptoms.
Fatigue
Fever
Chills
Cough
Sore throat
Runny or stuffy nose
muscle or body aches
Headaches
Headaches
ess commonly, vomiting and diarrhea
Monitor for secondary bacterial infections - sinus, ear, lung, pneumonia
If child is reporting fever sxs with no rise in temp ~
Concern for CDR1, innate immunodeficiency (mold/NK cell fxn)
Increased risk factor for autoimmune activity


81


82

## SARS-CoV-2

Multiple entry routes into the brain - olfactory bulb, thalamus, and brain stem may be infected through a trans-synaptic transfer of the virus. Additional vagal nerve delivery via dendritic cells.
Induces release of chemokines, cytokines, and inflammatory signals to the BBB and infects the astrocytes, which causes neuroinflammation and neuron death; neurodegenerative implications.

Pathogenic effect on the CNS with specific impact on the midbrain dopamine neurons which abundantly express ACE-2 receptors,
Spike protein can reach different brain regions, irrespective of viral brain replication. Can itself cause BBB dysfunction and damage neurons either directly, or via activation of brain mast cells and microglia and the release of various neuroinflammatory molecules.
Spike protein alters microglial purinergic signaling in vitro, may potentiate the Cell Danger Response
Published case report examined adolescents who acutely developed new OCD, neuropsychiatric, and motor dysfunction symptoms consistent with PANS, having a temporal correlation, 2 weeks after a diagnosis of Covid-19.
"Highly likely that neural autoantibody production is facilitated by SARS-CoV-2 infection..."



## Cytokine cascade

Entry in the brain via ACE2 (abundantly expressed in midbrain.)
TLR or NF-KB signaling activate the proinflammatory self-defense inflammasome after viral inflammatory
Pro-inflammatory feedback loop activates CNS immune cells, astrocytes and microglia, which induce IL-1, LL- 6 , TNF- $\alpha$, and IL-8.

Several CNS-related illnesses are linked with elevated levels of these inflammatory cytokines.


85

## Periodontal infections

A major under-recognized contributor to PANDAS/PANS and neuroinflammation. ID via qPCR Next-Generation Sequencing. Also tests for resistance in strains.

Dentist or periodontist collects a small amount of fluid from an infected pocket in the gums, as well as a saliva sample.
Avoid ozone for 1 week prior to sample collection as it's a potent antimicrobial.
Treatments using ozone are well-tolerated by P/P ~
Multiple published case studies using ozone gas to treat "untreatable" periodontal conditions ( $3-4$ month nightly rinse.)
Some evidence (15-day trial) ozone rinse is not as effective against gingivitis as commonly used chemicals (chlorhexidine) but is a viable alternative for chemicallysensitive. Need a longer duration study - empirically quite effective.

PMD: 3657058, 32594645



89

## No human effects?

Affects shikimate pathway - not found in human cells but is utilized by our gut microbiome
Reduces gut immunity and confuses the definition of "self", increasing the incidence of autoimmunity via Th17 and mast cell infiltration
Salmonella and Clostridia are resistant to it
Glyphosate-induced intestinal dysbiosis impacts CNS, in emotional, neurological and neurodegenerative disorders
In mice, low-level "subchronic" exposure increased anxiety and depressive-like behaviors
Low-level exposure linked to gut dysplasia
Animal studies, low-level maternal glyphosate exposure linked to increased incidence of ASD
PMID: 31422459, 29635013, 20012598, 28848410, 32398374

## Glyphosate (Roundup)

Genetically modify crops to be "roundup ready".
Allows the GM plant to survive the mechanism of the chemical.
But not just for killing weeds anymore!
Additionally used as a desiccant for non-GMO grains, spraying enough to kill the greenery via desiccation for easier harvest of grains, equating to higher than approved levels just before harvest.
Increases incidence of Fusarium mold infestation in storage.
"Coherent and compelling evidence that glyphosate and glyphosate-based formulations are a cause of non-Hodgkin lymphoma (NHL) in humans exposed to these agents."
Successful legal case linking exposure to NHL resulted in it being quietly taken off the market for residential use.
Commercial use allowed to continue!
PMD: 34052777, 31442895
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## Glycine backbone



Gly-phosate


Core of the molecule is glycine $\therefore$ may displace glycine metabolically

Impacts ~
Neurotransmitter (calming NT)
Glutathione (one of the AAs)
Glycine membrane channels (leading to channelopathies)
Possible fertility impacts ~ alters testicular morphology and testosterone levels

Glyphosate urine test


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## Atrazine - "pre-emergent" herbicide

Endocrine disrupting chemical with neuroendocrine/epigenetic toxicity.
Targets hypothalamus-pituitary-gonadal (HPG) axis.
Frogs: low exposure males become females, high exposure males can procreate.
Evidence of crosstalk between systems affected by Atrazine exposure, causing widespread dysfunction and leading to changes in behavior, even with no direct link to the hypothalamus.

EU banned Atrazine use in 2003 recognizing the health risks of Atrazine exposure as a public health concern with no way to contain contamination of drinking water.

Yet, the US recently reapproved Atrazine's use in the fall of 2020.
PMD: 27413107, 35410624
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## MOLD

Natural function of fungi is to compost and recycle
Excrete $1^{\circ}$ and $2^{\circ}$ metabolites ~ inhaled, ingested, and dermally absorbed
$1^{\circ}$ metabolites $\sim$ necessary for survival aldehydes, alcohols, odors, digestive enzymes, and structural elements (ie: beta-glucans)
$2^{\circ}$ metabolites ~ competitive antimicrobials, mycotoxins (energetically expensive for the mold to make)


Atrazine urine test

| Organophosphate pesticides |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Test Name ( $\mathrm{mcg} / \mathrm{g}$ ) | In Control | Moderate | High | Current Level |  |
| Diethyldithiophosphate (DEDTP) | 50.20 | 0.21-0.48 | 20.49 | 0.02 | 4.19 |
| Dimethyldithiophosphate (DMDTP) | $\leq 0.80$ | 0.81-5.08 | 25.09 | 0.29 | 5.75 |
| Diethylthiophosphate (DETP) | $\leq 0.70$ | 0.71-2.76 | 22.77 | 0.17 | 7.49 |
| Dimethylphosphate (DMP) | \$5.20 | 5.21-37.19 | 237.20 | 0.19 | 3.11 |
| Diethylphosphate (DEP) | 50.80 | 0.81~12.59 | 212.60 | 0.76 | 3.50 |
| Dimethylthionhosphate (DMTP) | $\leq 460$ | $461 \sim 2920$ | 32921 | 420 | 982 |
| Atrazine | $\leq 0.02$ | 0.03-0.05 | 20.06 | $<0.01$ | 7.16 |
| Atraine mercapturate | <0.03 | 0.04-0.06 | 20.07 | 0.03 | 7.04 |

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98



## Multisystem Multisymptom



More common than not that each
person in an exposure environment
has a completely different
presentation.
Every living being is affected.
Depends on type of mould,
presence of mycotoxins, duration
and dose of exposure, and
individual susceptibility.

[^0]Comparative Study >JAssoc off Anal Chem. 1983 Novi;66(6):1485-99.
© 2023 D r Jill Crista
Analysis for Fusarium toxins in various samples implicated in biological warfare in Southeast Asia
J Mirocha, RA Pawlosky, K Chaterejee, S Watson, w Hayes
PMID: 6643363
Abstract
Samples of leaves, water, cereal grains, soil, and yellow powder as well as blood, urine, and body



occur naturally on the substrates described and were correlated with the so-called "yellow rain
chemical attacks against tmong people in Southeast Asia Analysis of leaves, soi, water, and
chemical atactack against Hmong people in Southeast Asia. Analysis of leaves, soi, water, and
staged did not contain any Fusaium toxins. Mreoverer, T-2 and $H T-2$ thexins were found in buman


Mycotoxins have a long history of use as a biological warfare weapon.
"Yellow rain" T-2 toxin use against the Hmong people in Southeast Asia. So, yes, they affect everyone



109



Gather your Category Scores from the 3 previous categories.
CATEGORY I SCORE:
CATEGORY 2 SCORE: $\qquad$
CATEGORY 3 SCORE:
alculate your total mold risk:
TOTAL MOLD RISK
-4 -
10+ = Probable Mold- or Biotoxin-Related Illiness

THERTHINGSTO CONSIDER:
.
Inessinal parasites
Chronic viral sydid
Fon sensisivies
CVISS or inmunodeficienc s sndromes


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Mold Assessments


Direct Tests ~
Urine mycotoxin (LC-MS method)
Stool microbial assay +yeasts
Indirect Tests ~
Visual Contrast Sensitivity (vcstest.com)
Serum IgE/G *mycotoxin* antibody
$\mathrm{lg} E / \mathrm{G}$ mold spore antibody (standard) IgE/G mold spore antibody (stand
Urine mycotoxin (ELISA method) Urine
$\downarrow$ WBC, relative lymphopenia
$\uparrow$ NLR, microcytic anemia
Vit D ( $\downarrow 25-\mathrm{OH}$ and $\uparrow 1,25$ )
Liver function - esp $\uparrow G G T$
$\downarrow$ NK cell *function* with $\downarrow$ or normal NK cell total
†MMP-9 (mast cell correlate)
Organic Acids Urine Test
NeuroQuant (1' neuro sxs)
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## Voltage-gated channelopathies

Synaptic vesicles in the presynaptic terminal contain a high density of voltage-gated Ca2+ channels

When an action potential arrives at the distal end of the axon - the presynaptic terminals - the inrush of positive charge activates voltagesensitive $\mathrm{Ca} 2+$ channels.

Ca2+ entry then initiates the release of NTs into the synaptic cleft.
When NTs, such as ACh or glutamate, activate cation (for example $\mathrm{Na}+$ or $\mathrm{Ca} 2+$ ) channels, and are thus depolarizing, they can be described as excitatory.
PMD. 15753022
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Screen reliance


Only safe space for many P/P kids is virtual spaces
Virtual school often required
Screen addiction common (dopamine)
eEMF Blocking ~
Lap pads, sleep canopies
eEMF Grounding ~
Nature!, grounding mats
eEMF Discharging ~
Movement (produces non-polarized internal or iEMFs)


121

## Mercury sources

Organic ~
Methylmercury - fish/water contamination from coal-fired power plants
Ethylmercury (Thimerosal) - vaccine adjuvant, preservative (ie: contact lens solutions)
Inorganic/elemental ~
-"Silver" dental amalgams (about 50\% mercury).
-Dentists like its malleability and hardness as compared to other materials.
-Continuously release elemental mercury vapor.
-Amalgam surface area that exceeds the safe level of airborne mercury in the intraoral cavity:

Adult: >0.8 surface of a tooth
Child: >0.6 surface of a tooth
$-\therefore$ more than one small filling is harmful to a child's health
PMD: 21782213, 34941760

## Mercury

WHO March 2017 ~

- Considered by WHO as 1 of the top 10 chemicals or groups of chemicals of major public health concern.
Even small exposure amounts may cause serious health problems,
and is a threat to the development of the child in utero and early in life.
- May have toxic effects on the nervous, digestive and immune systems, and on lungs, kidneys, skin and eyes.
Comparison of typical exposures versus regulatory safety standards suggests that many people receive unsafe exposures.
1 in 6 women have mercury levels high enough to create a neurologic risk for their children.
Norway was the first country that banned the use of mercury in all products in 2008 including dental amalgam, followed by Sweden and Denmark.
2018: the EU banned the use of dental amalgam for children under 15 and for pregnant/nursing women.
PMD: 34941760, 24420334
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## Dental mercury amalgams in children

Evidence of safety of dental mercury amalgams in children has been based on 2 key studies from 2006 known as the Children's Amalgam Trials; followed >500 children each over $5 / 7$ years.
Both studies found no difference in neurobehavioral outcomes between the amalgam group and the composite (non-amalgam) group - although in both trials the amalgam group showed a statistically significant increase in urinary mercury levels.
These two studies, in addition to being widely cited in the literature, are cited by the FDA and the ADA as providing evidence for the safety of amalgam.
However, a 2011 reanalysis suggests harm, and >boys with common genetic variants Reanalysis used an exposure metric based on amalgam size and years of exposure -Found a significant association between amalgam and the porphyrin biomarkers for mercury-related enzyme blockage
"Dental amalgams are a significant
chronic contributor to mercury body burden."
PMD: 24420334, 21053054

## Exposure estimates

Organic mercury is more genotoxic than inorganic/elemental, yet "Amalgam-related
Hg exposure [which is inorganic/elemental form] exceeds that from fish or other sources for the majority of the population."
The highest allowable average mercury concentration in fish per serving when eating 1 serving per week $=0.46 \mu \mathrm{~g} / \mathrm{g}$

Whereas, estimates of Hg exposure from amalgam fillings "based on the least conservative of the scenarios evaluated, it was estimated that some 67.2 million Americans would exceed the Hg dose associated with the reference exposure leve (REL) of $0.3 \mu \mathrm{~g} /$ cubic meter of air established by the EPA."
Exposure estimates are consistent with previous estimates presented by Health Canada, and amount to 0.2-0.4 $\mu \mathrm{g} /$ day per amalgam-filled tooth surface, or 0.5-1 $\mu \mathrm{g} /$ day per amalgam-filled tooth, depending on age/other factors.
PMID: 21782213, 34941760

## Dental amalgams disperse

Mercury doesn't stay in the tooth!
A study quantifying the excretion and distribution of mercury in biological samples after dental -Concentrations of Hg in the biological samples of those with amalgams were found $6-8$ time higher than the non-amalgam users (control).
-Spike in Hg in RBCs, plasma, and urine on 1st day of filling, but not in hair or nails.

- -Accumulation in hair and nails by day 12 , but reduced in RBCs, plasma, and urine.

Mercury levels in the blood, urine or other biomarkers do not reflect the mercury load in critica organs.
Gestational mercury exposure ~
-Gestational exposure in infants of mothers who did not consume fish, had an elevated risk of URIs requiring a doctor visit.
-Alterations in both T cells and gene expression in placenta at birth.
Amalgams continuously release elemental mercury vapor (up to 20 micrograms per day.) Odorless and tasteless.
Primarily absorbed in lungs where it can disperse widely, even xBBB.
PMD. 2746468030773244.34129899

## Dentists and dental hygienists

Study of dentists in Iran found that the mean of the mercury level in the urine, nail, and blood was higher than the standard of the WHO.
"So, in accordance with Article 10 of the European Union Regulations (EUR), in the context of the Minamata Convention (MC) on Dental Amalgam (DA), in order to avoid the dangers of mercury exposure in dentists, it is necessary for Iran and other countries to approve laws and to implement a national plan to reduce mercury levels and replace the appropriate materials."
"Numerous studies have reported neurobehavioural effects in dental personne occupationally exposed to chronic low levels of mercury (Hg)."

- elevation of amyloid protein expression, deterioration of microtubules and increase or inhibition of transmitter release at motor nerve terminal endings.
- neurodegenerative diseases such as Alzheimer's, MS and mood disorders.
- idiopathic disturbances in motor functions, cognitive skills and affective reactions.

PMID: 33312669, 30589214

## Amalgams vs fish

Reference Dose of safety - level of exposure that is reasonably certain to be without appreciable risk for a population exposed over a long period of time.

EPA set RfD for methylmercury consumption in women of childbearing age (and their fetuses). No other population of defined, not even children
Reference Dose $=0.1 \mathrm{mcg} / \mathrm{kg} /$ day methylmercury. [45 lb child $=2 \mathrm{mcg} /$ day $]$
Amount of elemental mercury vapor from one amalgam filling $=$
1 surface = up to $20 \mathrm{mcg} /$ day.
The lower the body weight, the more increased the concentration.
"Throughout the world, efforts are underway to phase down or eliminate the use of mercury dental amalgam." (PMD 24420334) Yet there are no RfDs set for amalgams in the US, not even for those with lower body weight.

I acknowledge this is comparing different forms of mercury and so may have different health/absorption/accumulation effects

## Mercury health impacts

Neuro ~
As vapor: can $\times$ BBB and lipid cell membranes, and can be accumulated into the cells in its norganic forms.
Methylmercury can xBBB and placental barriers, causing serious damage in the CNS
Animal studies: motor and cognitive impairment and neural loss
Nephrotoxic.
Oral microbiome
Marked differences in the composition of the oral microbiome, associated with dental decay, found with even low concentrations of salivary mercury.
Gut ~
Gut connection to neurotoxicity: Healthy intestinal microbiota demethylates MeHg and promotes excretion through feces.
But in so doing, it impacts the gut microbiota and metabolites related to gut-brain interactions Induces changes of intestinal microbial community structure which induces changes to regulating neuron activity
Elemental Hg induces archaea (methanogens) conversion to methylmercury in vitro.
M10: 29777524, 32887894, 391818252, ,3324208
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129

## Mercury, mast cells, and histamine

Mercury induces histamine release from basophils.
Mercury induces inflammatory mediator release from mast cells, specifically VEGF and IL-6.
Animal models:
Induces a Th-2-dominated autoimmune syndrome with tissue injury in the form of a vasculitis and arthritis.
Sensitizes mast cells for mediator release and interleukin-4 expression.
Impacts mast cell survival.
Links to autoimmunity, disruption to BBB and subsequent neuroinflammation.
PMD: 20222982, 11222498, 19604304, 22103852, 9492216

## Mercury s/sxs

Symptoms are variable and nonspecific. Neuropsych sxs have high cross-over with P/P. Poor resistance to infection, especially to yeast and yeast overgrowths.
Anxiety, depression, "mercurial mood", irritability, suspicious, impulsive
Memory problems, incoordination, movement abnormalities, a sense of internal vibration, paresthesias particularly of the hands and feet
Neuromuscular junction: fasciculations, tremors
Halitosis, excessive salivation, metallic or salty taste, aphthous ulcers, tongue or tooth shocks, sensitive teeth, frequent dental caries, gingivitis, gums bleed easily, burning mouth syndrome, acute or chronic pharyngitis, perioral rashes
Night sweats, over-sensitivity to changes in temperature (think thermometer) and environments Intestinal Methanogen Overgrowth (IMO), IBS
Urinary frequency, kidney conditions
Increased allergic and mast cell related conditions
Increased fasting blood glucose
Increased risk for autoimmune conditions, esp Hashimoto's
"Tip: for toxic metals, look up homeopathic materia medica for complete list of Sx

## Mercury diagnostics

```
Due to rapid dispersal from plasma into tissue, and tissue accumulation, blood reference ranges are often not reliable indicators of health impact
Blood~
May apply to organic sources (diet and injected)
If positive, consider active/very recent exposure
\(>1.8 \mathrm{mcg} / \mathrm{L}\) associated with risk of Thyroglobulin Ab (Hashimoto's)
Urine ~
May apply to elemental sources (amalgam vapor)
Both pre- and post-provoked may be helpful
Why provoke? Assess chelating agent efficacy
Elevated unprovoked urine levels (95th percentile) considered significant for exposure.
*Abstain from eating fish/taking fish oil supplements for 1 wk prior to sample collection
Correlated to higher levels of fasting glucose.
Empirically, also correlated to chronic Candida/yeast burden.
\begin{tabular}{|c|c|c|c|c|c|}
\hline \multicolumn{6}{|c|}{Xc wials} \\
\hline & &  & Recter & wreteeer & Oursoenerramee \\
\hline Aluminum & (A) & 1.6 & <15 & - & \\
\hline Antimony & (sb) & 0.074 & \(<0.18\) & & \\
\hline Assenic & (As) & 12 & <40 & - & \\
\hline Batium & (Ba) & 0.88 & <5 & - & \\
\hline Berylium & (Be) & <al & \(<0.10\) & & \\
\hline Bismuth & (Bi) & 0.091 & <0.8 & - & \\
\hline Caamium & (cd) & 0.35 & \(<0.6\) & & \\
\hline Cesium & (Cs) & 11 & \(<9\) & & \\
\hline Gadolinium & (6d) & <al & \(<0.5\) & & \\
\hline Lead & (Pb) & 2.1 & \(<1.1\) & & \\
\hline Mercury & (Hg) & 0.55 & \(<0.8\) & & \\
\hline Nickel & (Ni) & 7.7 & \(<4\) & & \\
\hline Paladium & (Pd) & <d & \(<0.2\) & & \\
\hline Platioum & (Pt) & <d & \(<0.1\) & & \\
\hline Tellurium & (Te) & <d & \(<0.2\) & & \\
\hline Thallium & (T) & 2.2 & <0.4 & & \\
\hline Thorium & (Th) & <al & \(<0.007\) & & \\
\hline Tin & (Sn) & 0.19 & \(<3\) & - & \\
\hline Tungsten & (w) & <d & <0.4 & & \\
\hline Uranium & (U) & <dl & <0.03 & & \\
\hline \multicolumn{6}{|c|}{urin creatinne} \\
\hline & & \(\underbrace{\substack{\text { Redit }}}_{\text {Ressul }}\) & \(\substack{\text { Referece } \\ \text { NTERNAL }}\) & & \\
\hline Creatinine & & 32.5 & \(35-240\) & & \\
\hline
\end{tabular}

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\section*{Pesticides (Insecticides)}

Notoriously persistent chemicals in the tissues of mammals, especially those higher up the food chain.

The dose makes the poison, or does it?
Chronic low-level exposure more detrimental than a single poisoning event.
Neurotoxins ~
The issue comes when the molecular target is shared by non-target species.
Critical need for improved translation from animal models to humans
Pesticides such as organophosphates are linked to increased risk of neurological dz and dysfunction in humans, including chronic organophosphate-induced
neuropsychiatric disorders in a time and dosage dependent manner
Easily absorbed: inhalation, any cutaneous/mucocutaneous, ingestion.

IIL PMD: 31197504, 21402100, 30144465, 35439576



\section*{Pesticides (Insecticides)}

Exert neurotoxicity primarily through the inhibition of acetylcholinesterase (AChE) Leads to a buildup of ACh in the synapse, and hyperstimulation of cholinergic receptors in the CNS/PNS. Acute poisoning "cholinergic crisis"
Dopaminergic neuronal cells ~
Significantly alter dopaminergic neurochemistry
Additive/synergistic effects of different pesticides that act on different targets within the dopaminergic system.
Promote severe ox stress, mainly due to mitochondrial dysfunction, accompanied by significant upregulation and activation of caspases, thereby leading to apoptosis.
n vivo: damaged brain mitochondria marked by significantly reduced levels of catalase, glutathione (GSH) and superoxide dismutase (SOD), and increased lipid peroxidation.

PMD: 31197504, 21402100, 30144465

Organophosphate use in the US


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Pesticide urine test
\begin{tabular}{|c|c|c|c|c|c|}
\hline Test Name ( \(\mathrm{mcg} / \mathrm{g}\) ) & In Control & Moderate & High & Current Level & Previous Level 102902024 \\
\hline Diethyldithiophosphate (DEDTP) & 50.20 & 0.21-0.48 & 20.49 & 0.02 & 4.19 \\
\hline Dimethyldithiophosphate (DMDTP) & \(\leq 0.80\) & 0.81-5.08 & 25.09 & 0.29 & 5.75 \\
\hline Diethylthiophosphate (DETP) & 50.70 & 0.71-2.76 & 22.77 & 0.17 & 7.49 \\
\hline Dimethylphosphate (DMP) & \$5.20 & 5.21-37.19 & 237.20 & 0.19 & 3.11 \\
\hline Diethylphosphate (DEP) & \$0.80 & 0.81~12.59 & 212.60 & 0.76 & 3.50 \\
\hline Dimethylthiophosphate (DMTP) & 54.60 & 4.61-29.20 & 229.21 & 4.20 & 9.82 \\
\hline Atrazine & S0.02 & 0.03-0.05 & 20.06 & <0.01 & 7.16 \\
\hline Atrazine mercapturate & ¢0.03 & 0.04-0.06 & 20.07 & 0.03 & 7.04 \\
\hline
\end{tabular}

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\section*{Vaccines}
A. Please allow me to preface this section with a humble admittance that exist in extreme uncertainty about this subject, and I am far from being an expert.

What I am is a "curious digger", a clinician researcher.
I'm presenting my understanding as of this moment, which is simply my own understanding.
I welcome scientific dialogue and hope we can, together, advance our collective understanding
I invite you to remain curious, and promise that I will too
\begin{tabular}{l} 
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\section*{Risk of prevention vs infection}

In my PANDAS/PANS patient population, vaccines have consistently induced flares, but obviously so have infections.
How to counsel parents on each of the \(72+\) vaccinations to fulfill the childhood vaccination schedule?
What is the risk ratio of infection:vaccination for each?
Problematic situation:
No data on PANDAS/PANS kids.
Even more problematic:
In fact, there's no placebo-controlled data in healthy children either!

Wait, . . . what!?

\section*{Vaccine safety}

NO single childhood vaccine or combination of childhood vaccines has been tested against non-vaccinated controls in clinical trials, ever

Only a few have been studied against "controls" but the "controls" were either ~
- the adjuvants, rather than inert placebo. (ie: PedvaxHIB) By design, adjuvants are intended to evoke an immune response.
- other experimental vaccines (ie: Pneumonia: compared against an experimental meningitis vaccine that has never been approved, and to this day is still not approved. Yet the vaccine was not only approved for licensure, it became the "control" for the next generation pneumococcal vaccine trial.)

CDC's own definition of placebo from their website ~
"Placebo: A substance or treatment that has no effect on living beings
usually used as a comparison to vaccine or medicine in clinical trials.
https://www.cdc.gov/vaccines/terms/glossary.htm|\#P
These "controls" do not meet the definition of placebo-controlled


\section*{More details on Aluminum and asthma study}

Exclusion criteria and stratification by eczema diagnosis:
From an initial population of 398,191 children, 15,036 (3.8\%) did not meet inclusion criteria, \(30,976(7.8 \%)\) had vaccine-related exclusions, and 25,188 (6.3\%) were excluded due to asthma diagnosed prior to age 24 months. The final study cohort comprised 326,991 children, among whom \(14,337(4.4 \%)\) were diagnosed with eczema before age 12 months
The incidence rate of asthma appeared to increase with increasing levels of aluminum exposure in the eczema and no eczema cohorts.
Among children with eczema after adjustment for covariates, cumulative vaccine-associated aluminum was positively associated with persistent asthma (adjusted hazard ratio [aHR] 1.26 per 1 mg increase in aluminum, \(95 \%\) CI 1.07, 1.49).

For children with eczema, the mean and median cumulative vaccine-associated aluminum were 4.07 mg (SD 0.60), and 4.18 mg (IQR 3.97, 4.43), respectively.
For children without eczema, the mean and median were 3.98 mg (SD 0.72 ) and 4.18 mg (IQR 3.93, 4.43), respectively.
PMD: 36100331

\section*{145}

\section*{Mercury controversy}

Claim: Methylmercury (found in fish) and ethylmercury (thimerosal as sodium ethylmercurithiosalicylate) have different health effects...?
They're both organic form. Studies?
2016 industry flyer - "Thimerosal use is still permitted in multi dose vaccines and contact lens solutions at concentrations of up to 100 and \(70 \mathrm{mg} / \mathrm{kg}\) respectively." *mg, not mcg

CDC's National Immunization Program statement in 2004 - "...thimerosal-free vaccine costs more than the thimerosal-containing vaccine-about \(\$ 12\) versus \(\$ 8.50\) per dose."
https://assets.thermofisher.com/TFS-Assets/CMD/Appication-Notes/AN-43141-ICP-MS-Mercury-Contact-Lens-Solution-AN43141-EN.pdf

\section*{Vaccine efficacy}

NO vaccination on the childhood schedule has been tested for efficacy against nonvaccinated controls in clinical trials - ever
Do childhood vaccinations actually protect from the disease they're targeted against? How could we know? It's never been tested in a clinical trial.
Duration of non-placebo "controlled" trials from which vaccines have been licensed track for an average of 4-5 days. (le: Hep B)

What about long-term health outcomes?
Human and animal data suggest vaccine adjuvants increase the risk of developing an autoimmune disease, including RA, SLE, Sjögren syndrome, autoimmune thyroiditis and antiphospholipid syndrome.

PMID: 26275795, 27477999
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\section*{Vaccine efficacy}

The full extent of inquiry is one retrospective analysis by Dr. Paul Thomas of over 11,000 children born into his pediatric practice, where he had a statistically significant cohort of his patients who's parents decided to forgo vaccination and those who vaccinated their children providing a treatment group and a control group
Among the vaccinated, \(25.16 \%\) had a family history of autoimmunity, whereas among the unvaccinated, \(31 \%\) had the same characteristic.
The Quality Assurance Analysis showed ~
Vaccinated children had 4-5x more health issues than the unvaccinated children,
including allergic conditions, asthma, neurodevelopmental conditions such as ADHD, and infectious diseases, including those for which they were vaccinated against.
*these results were using his vaccine-friendly plan, which staggers vaccinations to reduce immune aggravation and allow for clearing of adjuvants.
This paper was retracted and to this date there's been no discernible reason why
PMD: 33266457


150


\section*{Published Reanalysis}

Dr. Thomas's paper was retracted based on the complaint of one person-after it
had passed peer review, was published, and had been read by over 250,000 people.
"His complaint hinged on the supposition - unsupported by any data - that vaccinated children made their scheduled HCVs more regularly than unvaccinated implying that those unkept appointments led to fewer diagnose.
We show, here, new data from the same practice that the opposite is true.
We have shown, using a variety of exhaustive methods, that the anonymous reader's concerns that led to the retraction of Lyons-Weiler and Thomas (2020 were unfounded. ...we conclude that the paper was wrongfully retracted..."

SOURCE: James Lyons-Weiler PhD and Russell L. Baylock, MD. Revistitigg EXcoss Diagnoses of llnosses and Conditions in Chidran
Whose Parents Provided Ilformed Permission to vaccinate Them, Intemational Joumal of Vaccine Theory, Practice, and Research. September 26,2022 p. 603 https//ddi.iorg/10.56098/interva2. September 26, 2022 p. 603. https://doi.org/10.560988/ivppr.v2i2.
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153

Additionally, the comparison of the Highand Low-vaccinated patients aged 1,500 days or more shows that vaccine cessation leads to a reduction in many conditions (thus the increased relative risk in the vaccinated patients;[]""
1 study. 1 reanalysis.
Why hasn't this been repeated?

\section*{Does volunteer reporting work?}

VAERS put in place by the CDC to monitor vaccine safety; voluntary, complaints about time it takes to submit a report.

So HHS commissioned the Agency for Healthcare Research and Quality (AHRQ) for a pilot project with Harvard researchers to use machine learning to facilitate detection and clinician reporting of vaccine adverse events directly into VAERS. Goal: improve completeness, validity, and timeliness of physician-approved case reports to VAERS compared to the existing spontaneous reporting system.

EHR from all ambulatory care encounters in a large multi-specialty practice. Every patient receiving a vaccine was automatically identified, and for the next 30 days, their health care Dx codes, lab tests, and prescriptions were evaluated for values suggestive of an adverse event.
Protocol was reviewed in advance by the CDC's Clinical Immunization Safety Assessment (CISA) Network.
Electronic Support for Public Heath-Vaccine Adverse Event Reporting System (ESPP:VAERS) Report, P: Lazarus, 2010

\section*{Conclusion}
"Low reporting rates preclude or slow the identification of "problem" drugs and vaccines that endanger public health. New surveillance methods for drug and vaccine adverse effects are needed. Barriers to reporting include a lack of clinician awareness, uncertainty about when and what to report, as well as the burdens of reporting: reporting is not part of clinicians' usual workflow, takes time, and is duplicative. Proactive, spontaneous, automated adverse event reporting imbedded within EHRs and other information systems has the potential to speed the identification of problems with new drugs and more careful quantification of the risks of older drugs."

Response? CDC shut down the pilot project.
Researchers: "the necessary CDC contacts were no longer available and the CDC consultants responsible for receiving data were no longer responsive to our multiple requests to proceed with testing and evaluation."

Electronic Support for Public Heath-Vaccine Adverse Event Reporting System (ESP:VAERS) Report, P: Lazanss, 2010

\section*{Timing and ethics}

Once a vaccine is licensed, it is considered unethical to conduct a placebo-controlled clinical trial, as you'd be denying the child the standard of care . .
*even though licensure was granted without a placebo-controlled clinical trial.
We are in a pickle:
How do we weigh the risk of infection against vaccination with limited data in kids with neurological autoimmunity.
How do we stratify which child's immune system will go too far?

In fairness, many of the herbs and other natural treatments that I'll be teaching about have never been tested against placebo controls. I'm fully aware that natural is not the same thing as safe.

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157

\section*{Weighing the risks}

Doctor, did you know that because the CDC hasn't done the appropriate research, the determination of whether any vaccination is safe or effective for your patient, or whether it increases their risk for developing allergies, asthma, ADHD, autoimmunity, or even the very infection it's intended to treat, is being left up to you?
"We need more investment in vaccine safety science,"
Heidi Larson, Director, WHO Vaccine Confidence Project
Ultimately it's the parent's decision, with "informed-as-much-as-possible-with-limited data" consent.
Parents are beginning to sue doctors since they can't get compensation for injury or death from the pharmaceutical companies for anything classified as a "vaccine" (per 1986 The National Childhood Vaccine Injury Act.)

\section*{Did you know you were being asked to take on that much personal}
medicolegal risk?
https://aaronsiri.substack.com/p/clinical-trial-to-license-rotatea

\section*{Coverage}
 inform as much as possible.
Require parents to sign a consent form for either decision; vaccinate or not. Consider each vaccine separately.
Copy the entire package in
Federal Regulatory Code
-Pre-2006 approved: \(\S 201.80\) "The labeling shall be revised to include a warning as soon as there is reasonable
evidence of an association of a serious hazard with a drug; a causal relationship need not have been proved."
nttps://www.ectr.gov/currentttitle-21/chapter-//subchapter-C/part-201/subpart-C/section-201.80
-After 2006: § 314.70 "To add or strengthen a contraindication, warning, precaution, or adverse reaction for
which the evidence of a causal association satisfies the standard for inclusion in the labeling under \(\$ 201.57\) (c) of this chapter,"
https://www.ecfr.gov/currenttitte-21/chapter-//subchapter-D/part-314/subpart-B/section-314.70
Consent form includes, but not limited to ~
-The risk of the getting the infection (sxs, severity, duration)
" \(\underline{X}\) I have read the vaccine package insert for
\(X\) understand that this vaccine has not been compared for safety or efficacy against true placebo-control in dinical trials, nor has it been compared against non-vaccinated children in a clinical trial."
-Make sure parents are aware that therefore it's impossible to provide a true "informed consent".
Check your malpractice coverage for vaccine injury and injury from not vaccinating


\section*{Diagnostics}


Clinical diagnosis
PE and symptoms as clues

General diagnostics
Infectious triggers

Environmental triggers

\section*{III}

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\section*{Environmental triggers}

Top 7 from my clinical practice ~
1. Herbicides
2. Mold
3. EMFs
4. Mercury
5. Pesticides
6. Vaccine adjuvants
(Food dyes get a dis-honorable mention)
Commonality: all are neurotoxins and immunotoxins.
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\section*{Prognosis}

Do they grow out of it? In my practice, not without treatment
Seem better outwardly, as they learn how to cope better/not disturb others = extreme inward suffering with outward "norm'ing".
Some improvement even without treatment after the hormone swings of puberty calm down.
Most can get back to life but must prioritize health/minimize environmental and infectious exposures
With adequate treatment, most grow into independently-living adults with careers, hobbies, relationships, etc


166```


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